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Investigations in Molecular Recognition Element Selection and Single-Wall Carbon Nanotube Properties

Ryan Martin Williams
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Investigations in Molecular Recognition Element Selection and Single-Wall Carbon Nanotube Properties

Ryan Martin Williams

**A dissertation submitted to the School of Pharmacy at
West Virginia University in partial fulfillment of the requirements for the
degree of**

Doctor of Philosophy in Pharmaceutical and Pharmacological Sciences

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Abstract

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Ryan Martin Williams

The *in vitro* selection of Molecular Recognition Elements (MREs) is a powerful tool for identifying molecules useful in numerous applications. This is achieved using the iterative Systematic Evolution of Ligands by EXponential Enrichment (SELEX) process. This technique utilizes a large library of 10^9 - 10^{15} different molecules which is enriched for those that bind to a target of interest. It also can be designed to enrich for molecules which do not bind to other, closely-related negative targets. We have developed a novel variation of this process called Decoy-SELEX. This enriches molecules that bind to the target of interest, but focuses on what the MRE should not bind to by using highly stringent negative selections with multiple targets. We have used this novel technique to identify separate MREs for three targets. First, a single-chain Fragment variable (scFv) antibody fragment MRE was identified that binds to prostate cancer cells but not benign prostatic cells. This molecule will be useful in the specific detection and targeted therapy of prostate cancer. Then, a single-stranded DNA (ssDNA) MRE was identified that binds to the herbicide atrazine but not other closely-related molecules. This molecule will be useful in the rapid, field-use detection of environmental contamination by atrazine. Finally, a ssDNA MRE binding to the pesticide malathion but not its metabolites was obtained. This molecule will also be useful as an environmental sensor for malathion contamination.

Single-wall carbon nanotubes (SWCNT) are a relatively new class of materials with novel mechanical, electrical, and optical properties. These properties are dependent upon the chirality and length of individual tubes, and can be altered with modifications, including chemical conjugates, DNA-wrapping, or protein adsorption. Therefore, SWCNT have many potential uses, including incorporation into electrical or optical molecular sensors. In order to determine the future potential of DNA-wrapped SWCNT in sensing applications, chirality identification, and purification, we tested their effect on the polymerase chain reaction (PCR). We determined the reaction-limiting concentration for different DNA sequences and SWCNT chiralities in addition to determining that DNA can be PCR-amplified directly from a SWCNT scaffold. This study identified parameters important in future studies for SWCNT application. Additionally, we determined the effect of DNA-wrapped SWCNT on a green algae model organism compared to sodium cholate-wrapped SWCNT. We determined that there was no effect of either type of SWCNT on algal growth at the concentrations assayed. This is important as the manufacture and application of SWCNT increases, making the likelihood of environmental exposure more prominent.

Dedicated to

My parents, David and Jackie Williams, for shaping everything about
what I have and will become

My sister, Mary Grace Williams, for friendship and happiness

My grandparents and close family members for having more influence
than they know

My close friends for positivity and support

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CHAPTER 1

Introduction to *In Vitro* Selection of Molecular Recognition Elements

1.1 Introduction

The Selective Evolution of Ligands by EXponential Enrichment (SELEX) is a method of obtaining molecular recognition elements (MREs) that bind to a target of interest (Figure 1.1). This *in vitro* process is a powerful tool for selecting molecules useful in the specific detection or treatment of targets of interest. Selectivity for a certain molecule or disease state can be ensured in designing the experiment with expected use of the final product being considered.

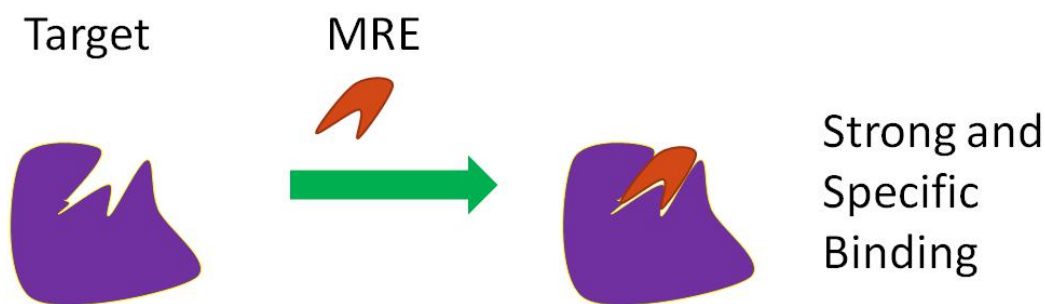


Figure 1.1. Visual depiction of Molecular Recognition Element (MRE) binding. A MRE is any molecule with strong and specific binding to a target of interest.

The SELEX, or *in vitro* selection, method obtains one or a few molecules that bind to the target with high affinity and specificity. These MREs are selected from a large library of 10^7 - 10^{15} random molecules. SELEX was originally described independently by the laboratories of Gold and Szostak in 1990 (Ellington and Szostak 1990; Tuerk and Gold 1990). These experiments selected RNA MREs that bind to proteins. Both ribonucleic acid (RNA) and deoxyribonucleic acid (DNA) MREs have been selected that recognize a range of targets, from single ions and small molecules to nanoparticles and proteins (Paborsky et al. 1993; Breaker and Joyce 1994; Huizenga and Szostak 1995; Gugliotti et al. 2004). These molecules are typically referred to in the literature as aptamers. The same SELEX method has been used to develop antibody fragment and peptide MREs that bind to small molecules, proteins, nucleic acids, and nanoparticles (Boder et al. 2000; Schaffitzel et al. 2001; El-Mousawi et al. 2003; Kase et al.

2004). The wide variety of MRE libraries and potential targets speaks to the numerous applications of SELEX.

The SELEX process uses a large library of random molecules that are incubated with the target of interest (Figure 1.2). Those molecules that bind to this target are kept and amplified, while those that don't are discarded. The amplified binding molecules are then incubated with negative targets, something the final MRE is designed to not bind to. Those molecules that do bind these targets are discarded and those that don't are amplified and re-subjected to the target of interest. This process of targeting and negative targeting is performed (rounds of selection) until one or a few high-affinity molecules remain in the library. The selected MREs are then

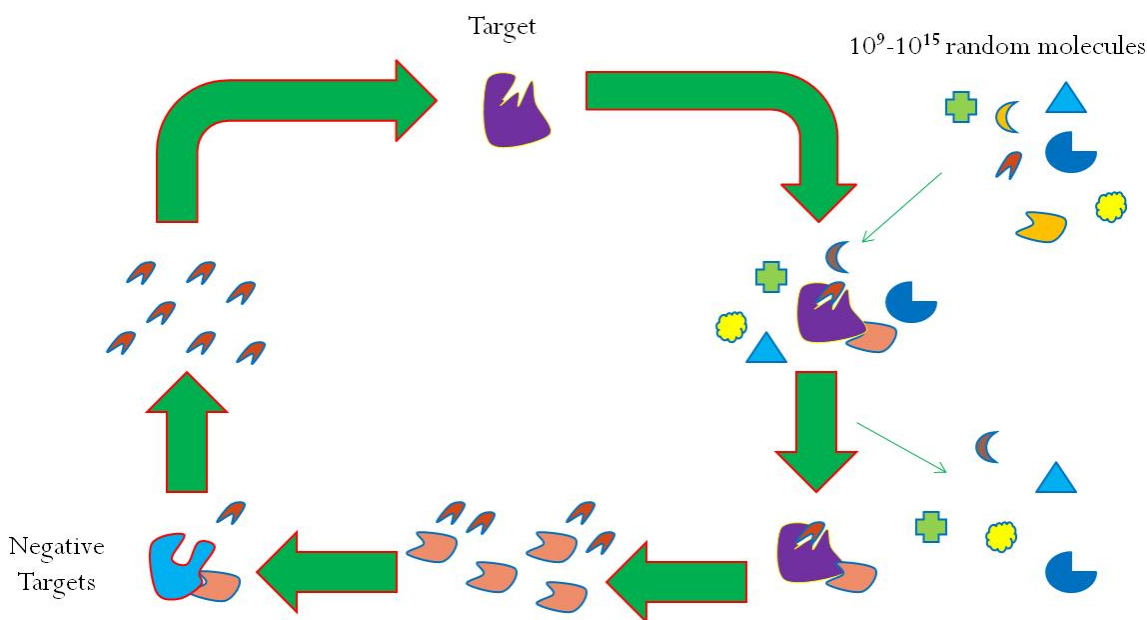


Figure 1.2. Scheme depicting the iterative SELEX process. A large pool of random molecules is incubated with the target of interest. Non-bound molecules are washed away while bound are amplified and subjected to negative targets. Non-bound molecules are amplified and re-subjected to the target under more stringent conditions.

assayed for their binding affinity and selectivity before they are used for various applications.

Regarding amino acid libraries, amplification typically occurs by use of a cellular host. For

DNA and RNA libraries, amplification occurs via the polymerase chain reaction (PCR).

Molecular recognition elements have been selected that bind to molecules displayed on the surface of cells. Originally performed by the Gold laboratory, DNA MREs that bind to the surface of red blood cells were selected (Morris et al. 1998). This work showed the ability to select MREs that bind to molecules displayed on cell surface membranes in their natural state. Since that time, SELEX has been used to select MREs that bind to cell surface molecules that are differentially expressed on the surface of cells. DNA ligands have been selected that bind to the vasculature of rat brain glioblastoma and not normal rat brain vasculature (Blank et al. 2001). Other nucleic acid MREs have been selected that bind to molecules on the surface of cancer cells but not other types of cells (Shangguan et al. 2008; Sefah et al. 2009; Zhao et al. 2009). This type of MRE selection provides the dual benefits of binding to anchored molecules and taking advantage of differences in surface expression of cells for the therapy and diagnosis of disease.

Compared to DNA and RNA MREs, amino acid-based MREs potentially provide the stability and specificity of antibodies *in vivo* (Table 1.1) (Nelson 2010; Li et al. 2011). Additionally, the non-reversible nature of their target interaction and high binding affinity is an advantage in tumor cell targeting. While having the positive attributes of antibodies *in vivo*, these MREs are selected *in vitro*. Therefore, more control over their binding target is possible for the experimenter. This includes the ability to preferentially select molecules which bind to a target but not a closely-related target. In the case of differential cell SELEX, differential surface molecule expression is taken advantage of for MRE binding (Cerchia et al. 2009). The result of amino acid MRE selection using differential cell SELEX is a molecule that has the advantages of antibodies with tunability in the experimental design.

Nucleic acid MREs have been investigated for their potential as *in vivo* diagnostic and therapeutic tools (Cerchia et al. 2002; Ireson and Kelland 2006; Lee et al. 2006; Guo et al. 2008;

Kim et al. 2009). It is possible that they will be degraded in the body by nucleases, however there are modification strategies to overcome this (Kanwar et al. 2011). In fact, Macugen (pegaptanib) is an anti-vascular endothelial growth factor (VEGF) nucleic acid MRE currently used in the clinic for macular degeneration (Moshfeghi and Puliafito 2005). This drug, however, is only used in the relatively closed environment of the eye, while Avastin (bevacizumab) is an anti-VEGF antibody used in anti-cancer therapy (Jain et al. 2006). Thus, there are modifications for using nucleic acid MREs *in vivo*, but amino acid MREs may be more useful.

Despite limited use *in vivo*, nucleic acid MREs have great potential for molecular detection in a variety of other settings. This is due to their typically strong affinity and selectivity, as well as their ability to be re-used. They can be denatured to release the target of interest after a binding event, and regenerated for further detection of the analyte (Ocana et al. 2012). Additionally, it is possible that nucleic acid MREs change their conformation upon binding their target, which can be capitalized on for detection transduction (Liu et al. 2010). Therefore, DNA and RNA MREs have a great deal of potential that is not found with antibodies or amino acid MREs.

There are advantages and disadvantages for DNA MREs compared to RNA MREs. Reports have shown that RNA MREs have a higher affinity for their target in general than DNA MREs (Potty et al. 2011). This is likely due to the inclusion of a 2' hydroxyl in the ribose backbone of RNA. However, nucleases that target RNA are much more abundant than those that target DNA, and thus identifying and applying RNA MREs is much more difficult. Additionally, PCR amplification of RNA is more difficult, as it must first be reverse-transcribed to DNA beforehand. Thus, DNA MREs typically are more stable and suitable for most applications, though RNA MREs do have value.

Discussed here is an overview on the creation of both amino acid and nucleic acid libraries for SELEX experiments. Then, selections for different types of each and finally, their applications, are investigated. Selection of antibody fragments for cell surface molecules and of nucleic acids for small molecules is focused upon.

Table 1.1. Summary of the advantages of the four most typical types of MRE libraries.

Type of Library	Library Subtype	Advantages
Amino Acid		- <i>In vivo</i> stability
		-Diversity of 20 amino acids,
	Peptide	-Well-studied -Widely available libraries
	Antibody Fragment	-Antibody structure and diversity selectable <i>in vitro</i> -Final product can be made into a full antibody -Pharmacological profile similar to antibodies which are clinically available
Nucleic Acid		-Large and diverse libraries easily obtained through combinatorial chemistry -Easily distributable (only need to know sequence to make your own) -Re-usable (can be removed from a target and regenerated for future use) -Easily modifiable
	DNA	-Easier to work with and less degradable than RNA -Very useful in molecular detection
	RNA	-Typically higher affinity than DNA -Useful for sensing

1.2 Methodology Overview

1.2.1 Amino Acid Library Creation

Amino acid libraries used in SELEX are typically displayed on the surface of living microorganisms. Generally, coding regions are inserted into phage, bacteria, or yeast that encode for surface expression of the randomized library. After expression of the library, it is incubated with the target cells of interest. Those that do not bind are discarded while those that do bind are kept and amplified. This amplification is performed by the natural replication of the microorganism. This enriched library is then incubated with the negative target cells. Those molecules that bind these cells are discarded while those that do not are amplified. Between every few rounds, the diversity of the enriched library is assayed by DNA sequencing of the amino acid library-encoding plasmid. The process continues until just one or a few specific and strongly-binding amino acid MREs are present. These are then assayed for their binding, specificity, and stability.

1.2.1.1 Phage Display

Bacteriophage have been used to display amino acid libraries. Libraries of short, random peptides that are displayed on the surface of phage have been used for SELEX experiments and are commercially available (e.g. Phage Display Peptide Library from New England Biolabs; Beverly, MA). These typically consist of 7-15 random peptides displayed by fusion to a surface protein. An example of this is the pIII protein of M13 phage, which is necessary for phage infection of *Escherichia coli* (Giebel et al. 1995). Random nucleic acid codons are synthesized and inserted into the plasmid for protein fusion. These libraries often consist of approximately 10^9 different peptides.

Additionally, a single chain Fragment variable (scFv) antibody fragment library has been constructed that is displayed on M13 phage (Pansri et al. 2009). An antibody fragment (Fab) library has also been constructed that is fused to the surface protein D of bacteriophage lambda (Huse et al. 1989). In this case, non-immune human antibody fragments are amplified from donors and constructed into a vector which displays the fusion protein on the virus surface. It has been noted that phage lambda is more capable of displaying large antibody fragments than M13 (Vaccaro et al. 2006).

Phage displayed amino acid libraries offer the advantages of rapid screening and amplification for the SELEX process. Selected phage are incubated with *E. coli* and plaques are produced. The replicated phage are then subjected to the next round of selection. The method used for SELEX is often panning (see below). While rapid, this method is not as efficient at isolating specific molecules as fluorescent activated cell sorting (FACS) (see below). A careful outline for the use of phage display has been published (Kay et al. 2001). Rapid and efficient amplification coupled with detailed protocols allow phage displayed amino acid libraries to be useful tools in cell SELEX.

1.2.1.2 Bacterial Display

Another platform for amino acid library display is through the use of bacterial hosts. A peptide display library of 12 random amino acids has been displayed as fusions to the *E. coli* flagellin gene (Lu et al. 1995). This library contains approximately 5×10^{10} different peptides and is available commercially (Invitrogen; Grand Island, NY). A library of the same size consisting of 15 random peptides has been displayed as a fusion to the *E. coli* outer membrane protein A (ompA) (Besette et al. 2004). These libraries offer the rapid and efficient screening necessary to create a MRE.

Antibody fragment display has also been performed in bacterial systems. A sub-library of scFv molecules in *E. coli* was created by PCR randomization of immune antibody fragments (Daugherty et al. 1998). Additionally, scFv molecules have been expressed as fusions to the surface protein A of gram-positive *Staphylococcus carnosus* (Gunneriusson et al. 1996). These constructs, however, have not been extensively explored for display of entire libraries or their use in SELEX experiments.

Bacterial display of amino acid libraries provides the benefit of rapid screening and amplification. Furthermore, they can be used in both panning and FACS-based separation (Kronqvist et al. 2008). While not as widely used as phage libraries, these libraries do not require multiple hosts, therefore increasing speed of the selection. In addition, detailed protocols for their use are available (Lu et al. 2003). These benefits render bacterial display of amino acid libraries a potential avenue for new developments in developing MREs.

1.2.1.3 Yeast display

Yeast display of amino acid libraries is a widely-used platform for generating MREs. A library consisting of approximately 10^7 random twelve amino acid peptides has been constructed (Krauland et al. 2007). It was formed as a C-terminal fusion to the Aga2p protein that is expressed on the surface of *Saccharomyces cerevisiae*. This library has proven to be of use in the selection of MREs.

The use of *S. cerevisiae* to display antibody fragments has also been extensively explored. This microorganism has been used to display a scFv library of approximately 10^9 different antibody fragments (Boder and Wittrup 1997). This library is freely available from the Pacific Northwest National Laboratory (PNNL; Richland, WA). A Fab sub-library has also been constructed that displays on the surface of yeast (van den Beucken et al. 2003).

Yeast display of amino acid libraries offers the advantages of use in FACS, as most include sequences for fluorescent protein tagging. Importantly, eukaryotic yeast have the ability to perform post-translational modifications not available in bacteria or phage. This allows for expressing the full diversity of antibody fragment libraries (Feldhaus and Siegel 2004). The described yeast systems also offer easy secretion and purification of selected MREs. Protocols for using these libraries are also easily available (Wang and Shusta 2005; Chao et al. 2006). These attributes allow the use of yeast in protein expression platforms to be an important tool in obtaining MREs by cell SELEX.

1.2.2 Nucleic Acid Library Creation

The design of a single stranded nucleic acid library for *in vitro* selection largely focuses on determining constant regions for PCR amplification. These are often designed by individual laboratories, however both DNA and RNA libraries are available for commercial purchase (e.g. TriLink BioTechnologies; San Diego, CA). A DNA library itself consists of two regions with known bases flanking a region with randomized bases. A RNA library also contains a promoter region to allow for reverse transcription of the library before PCR amplification. The greatest consideration in designing a DNA library is abrogating primer dimer formation and primer secondary structure effects. It is important to determine whether designed primer sequences will bind to each other or themselves (hetero- and homo-dimers), or if their reverse complement sequences will bind to one another, as all four sequences will be in the amplification reaction. Additionally, highly favorable secondary structures of primer sequences may inhibit PCR, and thus needs to be avoided for all four primer sequences. These can easily be interrogated using online PCR primer design software (e.g. Integrated DNA Technologies; Coralville, IA). The size of the randomized region must next be determined. This region must be designed to confer

the pool sufficient diversity, which can be represented by 4^n , with n being the number of random bases in the pool. For example, a pool with 34 random bases will have 4^{34} , or approximately 2.95×10^{20} , different possible sequences. Thus, longer random regions contain exponentially greater possible sequence space. In spite of this, longer random regions are more difficult to analyze for consensus binding sequences and are more likely to form secondary structures which inhibit PCR amplification. Therefore, random regions are typically between 20 and 80 bases, making overall libraries shorter than 150 bases (Hall et al. 2009). Longer regions are sometimes necessary, for example in obtaining catalytic MREs (Sabeti et al. 1997).

After design of the library for either DNA or RNA selections, it may be chemically synthesized using phosphoramidite chemistry (Hall et al. 2009). The library must then be purified from remnants of the synthesis reaction, and PCR amplification conditions optimized. During this optimization step, the library may not amplify properly, and it will be necessary to re-design it to suit the researcher's needs. Up to 10^{15} different sequences in the library are then incubated with the target of interest to begin the *in vitro* selection.

1.2.3 Amino Acid Separation Methods

A necessity of the SELEX process is separating molecules that bind to the target from those that don't. In cell SELEX, this is typically done by panning, magnetic sorting, or FACS. Often, a combination of these methods is used in order to most efficiently select MREs with high affinity (Table 1.2).

Panning, as used in cell SELEX, is a method of incubating the library with the target cells directly in the tissue culture well or flask. The unbound microorganism-displayed molecules are simply aspirated or decanted. The flask is washed typically three times taking care not to detach the target cells. The bound molecules are then amplified by replication of their host. This

process is most thoroughly described for phage-displayed libraries (Watters et al. 1997; McGuire et al. 2009) but has also been explored for yeast libraries (Wang and Shusta 2005). For phage, the bound molecules must be eluted and plated on bacteria for replication. In bacterial or yeast libraries, however, the appropriate microbiological media can be added to the cell culture flask or the human cells can be scraped from the flask. The enriched library-containing bacteria or yeast will amplify in this media, while any remaining human cells will not survive under these

Table 1.2. Typical methods of separation in SELEX.

Method	Commonly Used With	Advantages	Disadvantages
Panning	Amino acid libraries, some filter-immobilized targets with nucleic acid libraries	High throughput, quick	Less powerful separation, some undesired molecules remain after washes
Magnetic Separation/ MACS*	Typical for magnetic bead-immobilized targets with nucleic acid libraries, MACS can be used for amino acid library separation	High throughput	Limited space if using large cells (clogging), higher failure chance compared to panning with no advantages
FACS*	Typical for amino acid libraries	Very powerful separation ability with efficient tunability	Lower throughput, slower processing, expensive equipment
Non-magnetic target immobilization	Nucleic acid libraries; e.g. filter columns	Rapid, cheap, high throughput	Less powerful separation, may require many rounds
Capillary Electrophoresis	Nucleic acid libraries	Very powerful separation ability with efficient tunability	Expensive and expertly-used equipment

*MACS is Magnetic-Activated Cell Sorting. FACS is Fluorescence-Activated Cell Sorting.

conditions. Regarding negative selections, the unbound library molecules are removed by careful pipetting. The collected host organisms are then placed under optimal growth conditions. This method of separation is robust for large libraries. However, flocculation of the host

organisms or inefficient removal of non-binding molecules is possible. This creates an enriched library that contains some molecules that do not bind to the target. Therefore, panning is an efficient method of initial screening for the first few rounds of SELEX.

Magnetic sorting is another mechanism for separating binding for non-binding molecules. Often, magnetic-activated cell sorting (MACS) is used as a pre-enrichment procedure for similar reasons to those for panning. In the case of non-cellular targets such as proteins, the target is typically attached to magnetic beads and the library is passed through the column (Rice et al. 2006). A variation of this used a B cell antigen to bind chronic lymphocytic leukemia cells to magnetic beads for use with a phage library (Takahashi et al. 2003). The library molecules bound to the target are then eluted and replicated under appropriate conditions (or unbound molecules for negative selections). While useful for robust screening, this method may be difficult for use in cell SELEX due limited space within the column when considering human cells and library host cells are present. Therefore this method requires more preparation and greater potential for failure while performing the same pre-enrichment task as panning. It is for these reasons that the use of MACS for cell SELEX in obtaining amino acid MREs is rare in the literature.

Fluorescence-activated cell sorting (FACS) is often used to select MREs that bind to molecules on the surface of cancer cells. This method uses dual fluorescent labeling of the library and the target cells, which are removed from the culture flask. After incubation together, this mixture is subjected to FACS, wherein the software is instructed to collect events corresponding to double fluorescence. For negative targeting, collecting single fluorescent events corresponding to the library alone will render molecules that do not bind to the target. A basic protocol for yeast-displayed scFv library SELEX using FACS has been described

(Feldhaus et al. 2003). Additionally, a bacterial-displayed peptide library has been used for cell SELEX through the FACS method (Dane et al. 2006). The greatest advantage of FACS is the tunability of separation parameters and extremely efficient enrichment of binding molecules. It does, however, take considerably longer to screen very large numbers of cells. This is exacerbated when using two largely different cell sizes, as is the case with human and bacterial or yeast cells. Also, expensive equipment and expertise in operation is necessary. Therefore, FACS is more efficiently used after the library has been enriched and it is not necessary to use a large number of host cells to ensure full diversity is represented.

1.2.4 Nucleic Acid Separation Methods

Just as in selection of amino acid MREs, it is necessary to remove molecules that are not bound to the target of interest. Typically, this involves immobilization of the target of interest onto a substrate. There are separation techniques, such as capillary electrophoresis, that do not require this however (Mosing et al. 2005).

Immobilization of the target of interest is the most common form of nucleic acid MRE selection. This can be done with magnetic beads for manipulation in solution by a magnet, allowing library molecules not bound to the immobilized target to be simply washed away (Stoltenburg et al. 2005). This method has been successful for both protein and small molecule targets (Wang et al. 2009; Kim et al. 2010). Additionally, both protein and small molecule targets have been immobilized onto non-magnetic beads for column chromatography-based selection of MREs (Jackson et al. 2009). Protein targets can also be adsorbed onto nitrocellulose filter paper, however this is more difficult for small molecule targets (Ulrich et al. 1998; Sayer et al. 2002). A variation of this technique is to immobilize the target onto a microfluidic chamber

for rapid MRE identification (Lou et al. 2009; Ahn et al. 2011). Immobilization of the target typically occurs by conjugation through combinatorial chemistry approaches or adsorption. Other immobilization techniques have been developed, however they all ensure facile separation of bound from unbound molecules.

1.3 Amino Acid Molecular Recognition Elements for Cell Surface Targets

1.3.1 Peptide MREs

Molecular recognition elements selected from peptide libraries have been extensively explored. Especially from phage display libraries, the cell SELEX procedure has been used to select MREs that bind specifically to tumor cells. Peptides are short sequences generally less than 25 amino acids. Libraries with a very large diversity can be formed drawing from the inherent diversity of the 20 amino acids. A theoretical maximum for a 25 amino acid library is $\sim 10^{32}$ (20^{25}) or 1.28×10^9 for a seven amino acid library. However, typical construction techniques and reasonable working volumes typically limit library size to around 10^{10} molecules. Additionally, there is some bias in insertion of particular amino acids (Krumpe et al. 2006). A breakthrough in peptide library construction was the introduction of codon phosphoramidites (Glen Research; Sterling, VA), which ensures no premature stop codon or insertional biases are present. Compared to antibody fragment libraries, there may be less inherent structure within the MRE, but randomness is ensured.

Peptide MREs in cell SELEX have largely been selected through phage display methods. An example of this is the selection of peptide MREs that bound to the B-cell lymphoma line A20 (McGuire et al. 2006). This experiment did not use differential cell SELEX, but they were able to differentiate lymphoma cells from other normal white blood cells. In another study, primary chronic lymphocytic leukemia (CLL) cells were the targets of phage selection using a modified

MACS protocol (Takahashi et al. 2003). This study also did not use negative cell lines, however a panel of peptide MREs were selected that bound to different types of blood cells. A study performed that selected peptide MREs for non-small cell lung cancer (NSCLC) cell lines also did not use a negative target cell (Chang et al. 2009). One of the selected peptides, however, was successfully used for *in vivo* chemotherapeutic delivery studies. It is therefore possible to select useful peptides without a differential component of the selection.

The likelihood of selecting one or a few specific MREs that are fit for their expected use is increased if negative targets are used. One study selected for peptides that bound to the breast cancer cell line BT-474 but not the benign cell line MCF 10A (Shukla and Krag 2010). This study even did a negative selection against the tissue culture flask itself. This, however, seems to be inherent if negative selections are performed in the same type of flask as positive selections. The selected peptides did not bind to any of four benign cell lines that were used for binding studies. They bound to some, but not all, tumor cell lines studied and internalized into some. Another study with breast cancer cells selected for peptides that bound to HER positive SKBR-3 cancer cells but not HER negative MCF-10A benign cells (Abbineni et al. 2010). The selected peptides showed homology to other ligands for HER2, however the actual cell surface target molecule was not determined. Other studies targeting a follicular thyroid carcinoma cell line (Zitzmann et al. 2007) and separate colorectal tumor cell lines (Rasmussen et al. 2002; Kelly and Jones 2003) have been performed. Separate studies using the metastatic prostate cancer cell lines LNCaP (androgen sensitive) and PC-3 (androgen insensitive) selected peptides that showed no real homology between the reported sequence sets (Romanov et al. 2001; Jayanna et al. 2010). As these peptide sequence families did not converge, it can be concluded that they were not binding to the same cell surface molecule. Therefore it is likely these could be used together

to differentiate the two forms of prostate cancer and potentially define cell surface molecules present in one but not the other.

There also have been variations of the phage display peptide MRE selection process. A bacterial peptide display library has been used to select molecules that bound to breast adenocarcinoma cells ZR-75-1 but not benign cells (Dane et al. 2006). A variation on phage-displayed peptide panning has been performed in a microfluidic system for semi-automated selection of MREs (Wang et al. 2011). The above examples of using peptide libraries for selecting MREs express their potential in quick and efficient screening of random molecules to obtain diagnostically or therapeutically useful tools.

1.3.2 Antibody Fragment MREs

Antibody fragment MREs have also been selected that specifically bind to tumor cells. Antibody fragment libraries can be in the form of scFv, Fab, or others. These libraries typically have diversity on the order of 10^9 - 10^{10} molecules. This well exceeds the natural potential diversity of the human immune system, which has a maximum of approximately 10^7 different antibodies (based on heavy and light chain combinations of genetic diversity). Therefore, antibody-like MREs can be selected from libraries with much greater diversity than *in vivo* antibody production. It is likely that strong and specifically binding antibody fragments will be selected for cancer cells using differential cell SELEX.

Multiple studies have selected MREs that bind to tumor cells but not normal cells or other types of tumor cells. Two separate studies selected scFv MREs that bind to androgen insensitive prostate cancer cell lines. One targeted PC-3 cells and did negative selections with normal prostate cell lines (Liu et al. 2004). The other targeted the C4-2B cell line, which is derived from LNCaP cells that were used as a negative target (Conrad et al. 2009). The selected

antibody fragments were used for separate purposes, but the differences in selection design make it unlikely the MREs bind to the same surface molecule. Other studies have selected scFv MREs specific for melanoma cells (Kupsch et al. 1999), hepatocarcinoma (Yu et al. 2005), and *ex vivo* breast cancer tissue (Jakobsen et al. 2007). Additionally, Fab MREs have been selected that bind to T-cell malignancies (Willemsen et al. 2008) and ovarian carcinoma cells (Figini et al. 1998).

Antibody fragment MREs offer the specificity and pharmacological characteristics of monoclonal antibodies. Additionally however, they offer the selectivity and relative ease in production provided by the differential cell SELEX process. Their multiple potential uses therefore make their selection a strong method for cancer cell studies using SELEX.

1.4 Small Molecule Nucleic Acid Molecular Recognition Elements

1.4.1 DNA MREs

Molecular recognition elements in the form of DNA have been obtained for a wide variety of targets. These include ions of lead, mercury, and arsenic, among others (Kim et al. 2009; Chung et al. 2012). They also include protein targets such as vasopressin, protein kinase C, VEGF, and many others (Williams et al. 1997; Mallikaratchy et al. 2006; Hasegawa et al. 2008). A variation of this includes the selection of DNAzymes, which are selected to have enzymatic activity (Zhu et al. 2012). Another is the selection of double-stranded DNA MREs that bind to transcription factors (Sooter and Ellington 2004).

There have been a large number of MREs selected for small molecule targets for a variety of purposes. These include the identification of molecules such as ATP within cells (Huizenga and Szostak 1995), detection of pharmaceutical or illicit drugs such as theophylline and cocaine in biological contexts (Liu and Lu 2005; Chavez et al. 2010). DNA MREs have also been selected for environmental monitoring and contaminant detection purposes. Examples

of this include identifying MREs for pesticides such as acetamiprid or a variety of organophosphates (He et al. 2011; Wang et al. 2012). It has been shown that small molecules are specifically able to intercalate between bases of a DNA MRE in a sequence-dependent manner (Lin and Patel 1997). Additionally, normal antibody production against small molecule haptens require use of an immunogenic carrier, whereas *in vitro* selection does not, thus increasing specificity of the final binding element (Askonas et al. 1970).

DNA MREs offer a simplified and straightforward selection protocol, as discussed before. Additionally, they are stable as compared to RNA MREs, plus regenerative ability compared to antibodies. They are also very useful for molecular detection, as many references utilize them in this manner. Therefore, selection of DNA MREs offers an ideal molecular detector for detecting small molecules in complex environments.

1.4.2 RNA MREs

RNA MREs were historically the first to be identified (Ellington and Szostak 1990; Tuerk and Gold 1990). These molecules were originally identified to bind to protein targets, and many more examples have since been reported. There are also examples of the selection of ribozymes, which have catalytic activity often reliant on heavy metals (Wilson and Szostak 1999; Robertson et al. 2004).

There have also been numerous examples of RNA MREs for small molecule targets (Codrea et al. 2010). Similar to DNA, there have also been RNA MREs selected for ATP (Dieckmann et al. 1996). Additional examples include antibiotics such as chloramphenicol, tetracycline, and kanamycin, among many other types of targets (Burke et al. 1997; Berens et al. 2001; Goertz et al. 2004). Small molecule binding to RNA MREs is likely to occur similarly to DNA MREs.

RNA MREs typically have a high affinity and specificity for their target of interest. Additionally, these molecules offer many of the same molecular detection applications that DNA MREs do. They are, however, less straightforward to select and more likely to be degraded if used to detect their target in complex matrices. There are examples of modifications at their 2' hydroxyl which lower their degradation, however this often negates their advantage of having a higher binding affinity (Cui et al. 2004; de-los-Santos-Alvarez et al. 2007). It is therefore more straightforward with a higher probability of successful detector integration to select DNA MREs than RNA MREs.

1.5 Molecular Recognition Element Applications

1.5.1 Amino Acid Molecular Recognition Element Applications

1.5.1.1 Therapeutics

Amino acids in the form of monoclonal antibodies have been widely used as therapeutics (Brekke and Sandlie 2003). Well-known examples include trastuzumab (HerceptinTM: Genentech Inc.; San Francisco, CA) for HER2 positive breast cancer treatment and bevacizumab (AvastinTM: Genentech Inc) that binds to vascular endothelial growth factor (VEGF). These examples, among others, show the potential of amino acid-based therapeutic intervention.

Peptide therapeutics have been developed and used for a variety of diseases. Examples of these are to inhibit infection (Oyston et al. 2009) and inflammatory diseases (Vally et al. 2006). A review of therapeutic peptides obtained by phage display selection is available (Ladner et al. 2004). Examples of commercial peptide therapeutics include HumulinTM (Lilly; Indianapolis, IN) for diabetes treatment and leuprolide (LupronTM: Abbott Laboratories; Abbott Park, IL) for metastatic prostate cancer treatment. Peptides obtained by cell SELEX are therefore possible alternatives to antibodies for cancer treatment.

Antibody fragments have also been used therapeutically. As of 2009, approximately 90 scFvs had been patented in the U.S., with about 30 being specifically for cancer therapeutics (Pucca et al. 2011). The humanized h5G1.1-scFv specific for complement protein C5 (pexelizumab: Alexion Pharmaceuticals; Cheshire, CT) has undergone clinical trials for its ability to reduce myocardial infarction after coronary artery bypass surgery (Smith et al. 2011). This molecule was not selected by SELEX, but was engineered from a previously-selected monoclonal antibody (Evans et al. 1995). The results of clinical trials are mixed (Lin 2011), however the pharmacological profile of the antibody fragments does appear to be safe (Fitch et al. 1999). It is just one of 19 scFvs to reach clinical development as of 2010 (Nelson 2010). There have also been 22 Fabs to reach clinical trials, 3 of which have been approved (Nelson 2010). While some of these antibody fragments were engineered from monoclonal antibodies, it is clear that scFv treatments obtained from SELEX will be useful and more specific for cancer treatment.

Another scFv molecule that binds to the HER2 receptor on the surface of tumor cells has undergone clinical trials. The antibody fragment was produced similar to the previous one, however it was conjugated to exotoxin A (scFv-ETA) (Wels et al. 1992). The results of the trials proved safe, and the drug was moderately effective (von Minckwitz et al. 2005). This use of antibody fragments in tumor cell treatments shows the potential for specific drug delivery of various therapeutics.

The use of amino acid MREs is clearly a potentially fruitful field. The experimental tunability for selectivity and *in vivo* stability has been previously demonstrated. Recently, theranostics, or the dual use of single molecules for therapeutic and diagnostic purposes, has become popular (Goldenberg et al. 1990). There is clearly the possibility of using amino acid

MREs that have been conjugated to contrast materials as in the previous study. The use of MREs obtained by cell SELEX in therapeutics and/or diagnostics has only begun to be explored but represents great potential.

1.5.1.2 Disease Detection

Amino acid MREs for detection of disease is a potential clinical application of these molecules. For biomarker-based cancer detection, blood is typically extracted and subjected to immunoassays with monoclonal antibodies. There also lies the potential for magnetic, radio-, or fluorescent labeling of these antibodies for diagnostics. Antibodies used in these assays do not always bind to molecules that represent disease states. For example, anti-PSA antibody used in prostate cancer detection has a false-positive rate over 75% (Thompson et al. 2006). These uses make it important to obtain molecules that specifically detect malignant cells.

Initial research using peptide MREs obtained by SELEX have been promising. *Ex vivo* human and *in vivo* mice studies using colon cancer-specific peptides have been performed (Kelly et al. 2004). Using radiolabeled and fluorescent peptides, favorable pharmacological profiles and detection were observed. Similar results were found using an anti-prostate cancer peptide selected by phage display (Zitzmann et al. 2005). While not extensively explored for clinical use, peptide MREs may have great diagnostic potential.

Similar studies to those performed with peptide MREs have been done with antibody fragments (Goldenberg et al. 1990). Another potential use of antibody MREs is in *ex vivo* diagnostics such as immunoassays or microfluidic cell capture. Though using a monoclonal antibody, previous studies have shown the ability to capture circulating tumor cells with antibodies in a microfluidic device (Goldenberg et al. 1990). There is the obvious potential for

application of antibody fragment MREs to a similar device. These and other uses make the diagnostic potential of amino acid MREs extremely important.

1.5.1.3 Surface Molecule Elucidation and Discovery

Whether the selected MRE is nucleic or amino acid, it can be used to learn more information about its target cell. An important attribute of cell SELEX is that no *a priori* knowledge of the cell surface molecular target is necessary. It is also likely that cells in disease states will express surface molecules differently than normal cells. No previous knowledge of this differential expression is necessary however, because the differential cell SELEX process will take advantage of this fact in selecting MREs.

The cell SELEX process has been used to select MREs which aided in identification of their own surface ligand. DNA MREs have been selected that bound to tenascin C overexpressing cells and not unaltered cells (Daniels et al. 2003). As predicted, the MREs bound to tenascin C. Another study determined that ICAM-1, an intercellular adhesion molecule, was the target of scFv MREs selected on androgen-insensitive prostate cancer cells (Conrad et al. 2009). DNA MREs have also been used to discover new proteins expressed on the surface of dendritic cells (Berezovski et al. 2008). These works show that overexpression of a surface protein does allow MRE selection against them. Additionally, novel or confirmatory roles of previously described molecules can be determined. There also lies the potential for discovery of new cell surface molecules or biomarkers. This use of MREs is potentially very important for a fundamental understanding of disease proteomics.

1.5.2 Nucleic Acid Molecular Recognition Element Applications

Nucleic acid MREs have great potential in multiple settings. Predominant among these is analyte binding in biosensors. Both DNA and RNA MREs have been incorporated into sensors

(Cho et al. 2009). Typically, these come in the form of optical, electrochemical, or mass sensitive sensors, sometimes with the incorporation of various nanoparticles for signal transduction (Song et al. 2008). As noted previously, nucleic acid MREs function well in this setting because they can be obtained for most targets, can be modified and incorporated into devices, are relatively stable, often change shape upon ligand binding, and are reusable.

Nucleic acid MRE-based sensors have been developed utilizing many different modalities. For example, sensors for just ATP have been developed using DNA MREs integrated into many types of sensors, including those utilizing field-effect transistors (Goda and Miyahara 2012), fluorescent detection (Urata et al. 2007; Wang et al. 2008), quantum dot luminescence (Huang et al. 2010), surface-enhanced Raman scattering (Li et al. 2012), carbon nanotubes (Zhang et al. 2010), microbalances (Ozalp 2011), electrochemical transducers (Zuo et al. 2009), and many others. These encompass the three main detection modalities mentioned above, and give just a small swath of the large spectrum of methods possible with nucleic acid MREs.

In addition to these many detection methods, MRE-based sensors have been developed for many different targets. These include small molecules such as ATP as noted above, cocaine (Zhou et al. 2012), tetracycline (Kim et al. 2010), and many others. They also include proteins such as thrombin (Cho et al. 2008) and anthrax protective antigen (Cella et al. 2010), and insulin (Yoshida et al. 2009), among others. It is noteworthy that many of these detection mechanisms are modular in that a separate MRE can be used within the same design to detect new analytes.

In addition to detection, nucleic acid MREs have been investigated for various biomedical applications. As noted earlier, MacugenTM (pegaptanib) is in the clinic for macular degeneration treatment. Other MREs have been used for *in vivo* therapeutic delivery of drug-

loaded particles (Wu et al. 2010). They can be integrated into the membrane of lipid-based particles, or immobilized onto the surface of certain manufactured nanoparticles, each having a drug-release mechanism (Yu et al. 2011; Li et al. 2012; Sundaram et al. 2012). Furthermore, they have been complexed with positron emitters, fluorophores, radiolabels, or other transducing elements for *in vivo* imaging of molecular expression (Tavitian et al. 2009; Shi et al. 2010; Yu et al. 2011). In spite of these applications, they have not been as widely moved into clinical settings as antibodies, largely due to drawbacks previously discussed.

In total, nucleic acid MREs have a wide array of potential applications. They have been studied for their biomedical uses; however their properties allow them to be more useful in biosensors. Thus, nucleic acids MREs will be the focus of molecular detection for sensing applications.

1.6 Conclusions

Selection of MREs using differential cell SELEX represents a powerful method of differentiating between cell types. This process requires no prior knowledge of cell surface molecule expression and takes advantage of differential expression profiles. Peptide and antibody fragment libraries, the host on which the library is displayed, and the selection method each have advantages and situations in which they are useful. Expansion of host scaffolds has been performed, even using cereal crops (Stoger et al. 2000) and human HEK-293T cells (Ho and Pastan 2009) to display engineered scFv molecules. As more MREs are being developed, it is certain that their uses and clinical investigation will continue to expand.

Nucleic acid MREs have a many ideal applications, and are able to be obtained for a wide array of targets. They yield a powerful method of molecular binding and differentiation of molecules. As discussed, they are particularly useful in biosensors for detection of various

analytes. Increasingly more detection modalities are being developed and are only limited by creativity of their integration. Therefore, the continued selection and investigation of nucleic acid MREs is important for a variety of applications.

Therefore, the work discussed herein developed different types of MREs for different applications. In total, the SELEX process was applied to obtaining MREs for three different targets. Specifically, this work developed and utilized a novel variation called Decoy-SELEX which focuses on what the final MRE should not bind to. As in classical SELEX, the library is enriched for the target of interest, however many stringent negative selections are performed. Antibody fragment MREs were obtained that could bind to prostate cancer cells and differentiate between them and benign prostatic cells. These have potential in surface molecule discovery, cancer detection, and targeted therapy. Separately, DNA MREs were obtained for two small molecule pesticides, atrazine and malathion. These have potential use as binding molecules in biosensors for these environmental contaminants. In total, each MRE has strong affinity and discriminatory power, as well as multiple potential applications building upon this work.

CHAPTER 2

***In Vitro* Selection of Single-Stranded DNA Molecular Recognition Elements for the Herbicide Atrazine**

2.1 Introduction

Atrazine is a chlorotriazine herbicide used widely in crop production. An estimated 76.5 million pounds are used per year in the United States alone (USGS 2002). Its use is estimated to increase crop yields 4-6% (Ackerman 2007). The economic benefits of atrazine use have led to its continued application for many agricultural processes.

The widespread application of atrazine, however, leads to contamination of aquatic ecosystems and water supplies. Dependent upon rainfall, up to 20% of atrazine applied to crops may run off or be leached from soils (Graymore M 2001). Streams bordering croplands applying atrazine have been shown to be contaminated at levels up to 1000 µg/L. Additionally, some drinking water reservoirs in the U.S. have up to 88.4 µg/L. Atrazine has been detected in surface and ground waters in the state of West Virginia (Pasquarell and Boyer 1996; Ator and Ferrari 1997). Removal of atrazine in water treatment plants is possible, but varies widely throughout the US (Jiang H 2006). The US Environmental Protection Agency (EPA) Maximum Contaminant Level (MCL) is 3 µg/L for human exposure to atrazine (USEPA 2011). This level has been exceeded in many surface waters tested in West Virginia (Ferrari and Assessment 1997). Furthermore, the European Union has banned use of atrazine due to contamination levels above recommended limits (Sass and Colangelo 2006).

The recommended limit of exposure is based on the potential negative effects of atrazine. Atrazine exposure at environmentally relevant levels affects aquatic organisms; including feminization of male frogs (Hayes et al. 2002), salamander desiccation (Rohr and Palmer 2009), and endocrine disruption in fish (Suzawa and Ingraham 2008). Endocrine disruption is also a potential mechanism of atrazine action in humans and other mammals (Cooper et al. 2007; Fan et al. 2007; Albanito et al. 2008). It has been shown to interact with a steroidogenic transcription

factor and a growth factor receptor, causing signaling pathways to be upregulated. This may lead to hormonally-linked cancers and reproductive abnormalities. Additionally, atrazine may affect innate immune function and increase damage caused by other toxins such as arsenic (Tchounwou et al. 2001; Mizota and Ueda 2006). There is, however, evidence that contradicts an EPA finding that atrazine is not a potential carcinogen (Mills 1998). It is therefore important that food and water supplies be routinely monitored for atrazine levels.

The current methods of atrazine detection are either time- and labor-intensive or are not specific for the herbicide. The most common method is gas chromatography, which requires extensive sample preparation (Amistadi et al. 1997). Combinatory chromatographic methods have been used to determine the presence of atrazine in both serum and urine (Brzezicki et al. 2003; Panuwet et al. 2008). Antibody-based assays are also used, however the antibody used cannot differentiate between atrazine and closely-related molecules (Wortberg et al. 1996; Suri et al. 2009). It is therefore necessary to identify an easily-synthesized binding element with high affinity and selectivity. Utilizing Decoy-SELEX (Systematic Evolution of Ligands by Exponential Enrichment) an atrazine-specific Molecular Recognition Element (MRE) was identified with sub-nanomolar affinity.

Molecular Recognition Elements are antibody-like molecules that strongly and selectively bind to a target of interest. They are produced by the *in vitro* selection process initially described independently by the laboratories of Gold and Szostak in 1990 (Ellington and Szostak 1990; Tuerk and Gold 1990). MREs are selected from a large (10^9 - 10^{15}) library of single-stranded DNA or RNA. The library is initially incubated with the target of interest. Molecules that do not bind to the target are discarded and those that do bind are amplified. These are then subjected to a series of negative targets, after which molecules that *do not* bind

are amplified. This iterative process continues, utilizing increasingly stringent conditions, until a MRE that specifically binds to the target of interest, but not negative targets, is identified. The selected MRE may then be used for applications such as detection of the target molecule.

A single-stranded DNA (ssDNA) MRE with high affinity and specificity for atrazine was isolated using a novel Decoy-SELEX process. This SELEX variation enriches the library for the target of selection but focuses on stringent negative selections against undesirable binding targets. Those negative targets are chosen by chemical similarities, such as target metabolites, or their environmental proximity to which they are found with the target. The selection scheme for atrazine was designed to ensure the MRE would bind to neither atrazine metabolites diamino-chloro-triazine (DACT) or desethyl atrazine nor the closely-related herbicide simazine (Muller et al. 1997). It was also designed so that the MRE would not bind to propanil, 2,4-D acid, or malathion, which are pesticides found at similar levels and environments to atrazine (Kucklick and Bidleman 1994; Coupe et al. 2000; Purcell et al. 2001; McCarthy and Fuiman 2008). These selected molecules may then be incorporated into a sensing device, allowing for specific atrazine detection (e.g. (Soh et al. 2009; Yang et al. 2010; Lin et al. 2011)).

2.2 Methods

2.2.1 Selection of Atrazine-Specific MREs

In order to select an atrazine-specific MRE, the Decoy- SELEX process was used (Figure 2.1). A single stranded DNA (ssDNA) library consisting of 10^{15} different molecules was screened through 12 rounds of selection. The ssDNA library, termed RMW.N34, consists of two 23 base constant priming regions for polymerase chain reaction (PCR) amplification flanking a 34 base random region (synthesized by Eurofins MWG Operon; Huntsville, AL). From this library, molecules that have a sequence and tertiary structure that specifically binds to atrazine

were selected. Molecules in the library which bound to closely-related structures and other pesticides were removed (Figure 2.2).

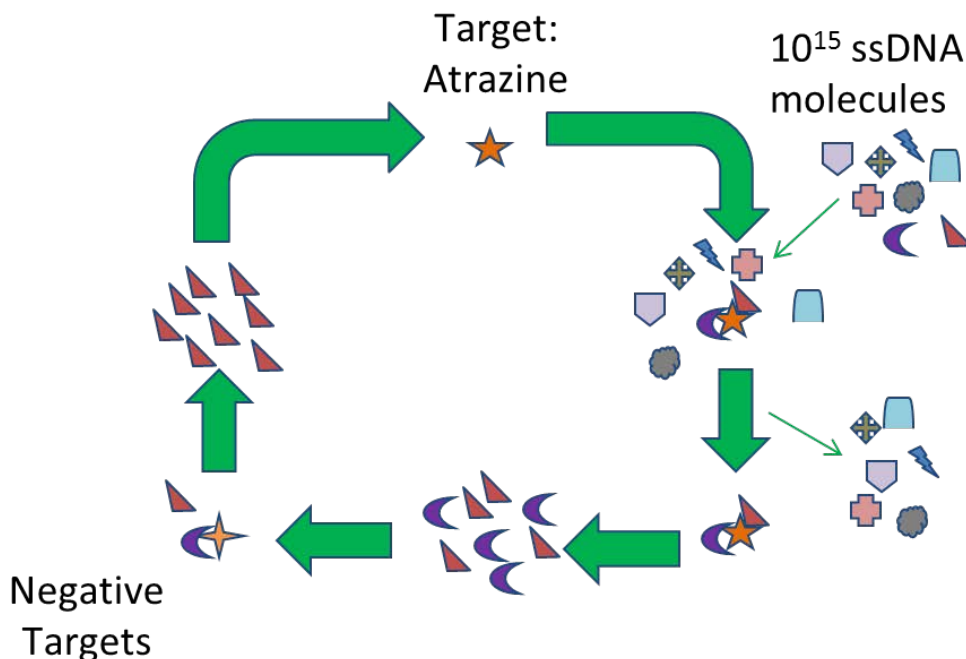


Figure 2.1. Scheme depicting the Decoy-SELEX process for atrazine MRE selection. A library of 10¹⁵ ssDNA molecules were incubated with the target atrazine. Those that bound were amplified and incubated with negative targets. Those that did not bind the negative targets were amplified and subjected to additional rounds of *in vitro* selection.

An atrazine derivative, desethyl atrazine (Figure 2b) (Fluka Analytical; St Louis, MO) was covalently biotinylated by Sulfo-NHS-LC-Biotin (Pierce; Rockford, IL) per manufacturer's instructions. Biotinylated desethyl atrazine was then incubated with streptavidin-coated magnetic beads (New England Biolabs; Ipswich, MA) to immobilize the selection target. Unbound desethyl atrazine was washed to leave only biotinylated desethyl atrazine bound to magnetic beads (immobilized target).

For positive immobilized target selection, the ssDNA library was incubated with 50 μ L immobilized target in 500 μ L of buffer consisting of 100 mM sodium chloride, 20 mM Tris-HCl, and 2 mM magnesium chloride (1X selection buffer; SB). After an incubation period with

rotation at room temperature, the immobilized target was separated from the solution by magnet, washed three times with 1 mL SB, and resuspended in SB. This suspension served as a template for PCR amplification of bound ssDNA. PCR conditions were as follows: enriched ssDNA library, 400 nM forward and biotinylated reverse RMW.N34 primers (Eurofins MWG Operon; Huntsville, AL) (forward: 5'-TGTACCGTCTGAGCGATTCGTAC-3', biotinylated reverse: 5'-Biotin-GCACTCCTTAACACTGACTGGCT-3'), 250 μ M deoxynucleotide triphosphates, 1X GoTaq Reaction Buffer (Promega; Madison, WI), 3.5 units *Taq*, and pure water. Reaction conditions were: initial denaturation of 95°C for 5 minutes; cycling at 95°C for 1 minute, 63°C for 45 seconds, and 72°C for 1 minute; and final extension of 72°C for 7 minutes. Large-scale amplification of 4 mL and 2 mL was done preceding positive and negative selection rounds, respectively. This selection scheme for the desethyl atrazine target occurred for Rounds 1-5 of selection, each time with a shorter incubation period.

Amplified dsDNA was then purified via the GFX PCR purification kit (GE Healthcare; Piscataway, NJ). Eluted dsDNA from the column was incubated with streptavidin agarose resin (Pierce; Rockford, IL). Having used a biotinylated primer for the reverse strand amplification allowed for binding of the amplified dsDNA to the agarose resin. This was placed into a flow-through column and washed with 5 volumes of 1X PBS. Five volumes of 1 M sodium hydroxide were then used to dissociate the DNA strands, eluting the forward strand ssDNA from the column. This solution was neutralized with 0.1 volume 3 M sodium acetate (pH 5.2), to which 2.5 volumes of cold 100% ethanol were added. Glycogen at 10 μ g/mL was added as a co-precipitate for ethanol precipitation, and the solution was frozen. After freezing, this was centrifuged at 13,000xg for 1 hour, washed with 70% ethanol, and centrifuged again for 15

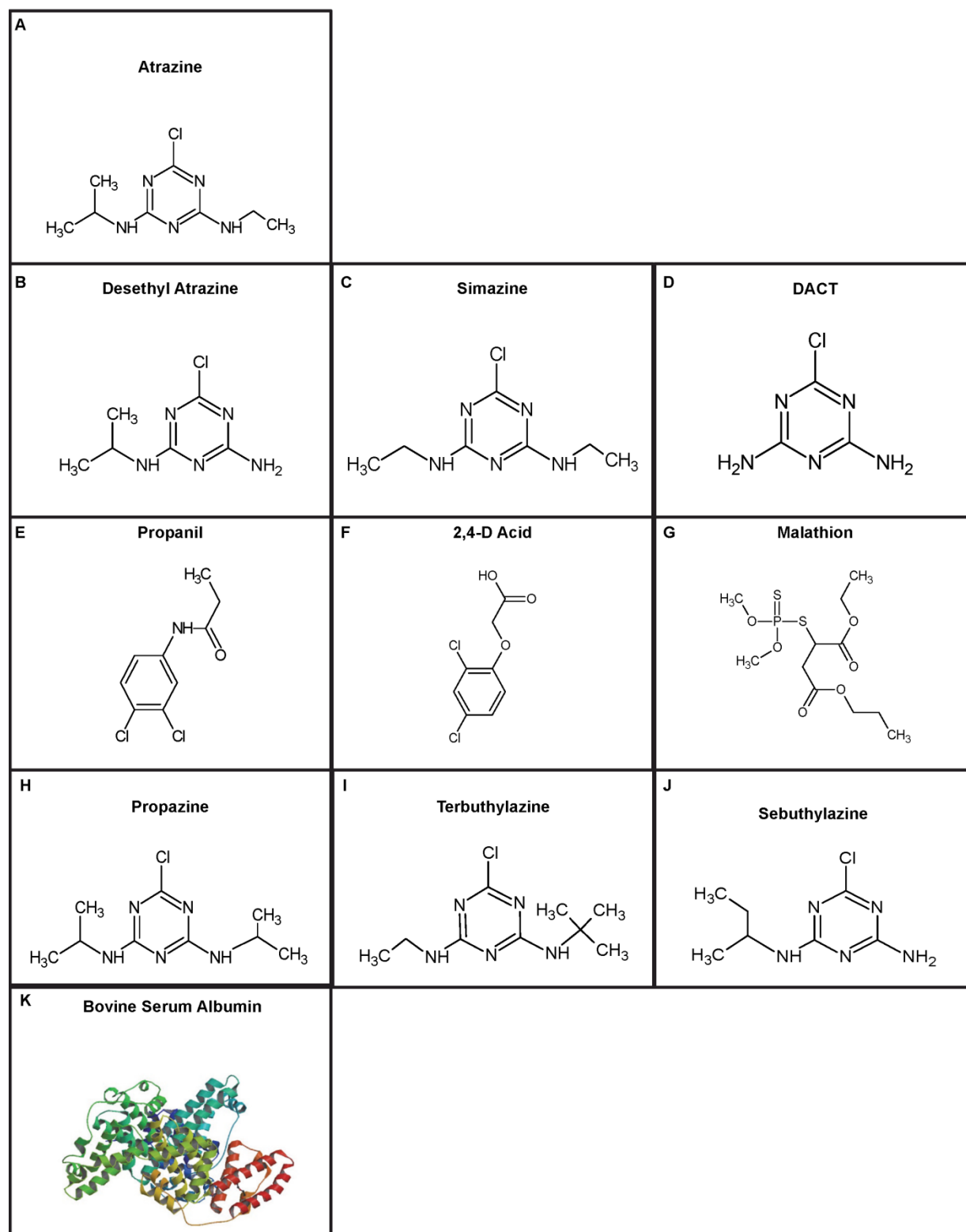


Figure 2.2. Chemical structures of molecules used in the Decoy-SELEX scheme and cross-binding assays. (a) Structure of the herbicide and target of selection atrazine. (b) Structure of atrazine derivative desethyl atrazine, used to immobilize onto magnetic beads. (c) and (d)

Structures of simazine and diamino-chloro-triazine (DACT): chemically similar to atrazine, and used in rounds of negative selection. (e), (f), and (g) Structures of propanil, 2,4-D acid, and Malathion: pesticides found in ecologically relevant environments to atrazine and used in negative selections. (h), (i), and (j) Structures of propazine, terbuthylazine, and sebuthylazine: structurally similar metabolites of atrazine used in cross-binding assays. (k) Ribbon structure of bovine serum albumin (BSA) used in cross-binding assay (Bujacz 2012).

minutes. The supernatant was then decanted and the ssDNA pellet dried under a vacuum. The purified ssDNA pellet was resuspended in SB and subjected to another selection incubation.

Negative selections were done to ensure the ssDNA MRE is specific for atrazine. Sulfo-NHS-LC-Biotin (Pierce; Rockford, IL) was blocked with Tris-HCl. Then the blocked biotin reagent was incubated with streptavidin coated magnetic beads and washed with SB (immobilization substrate). 50 μ L of blocked biotin reagent immobilized onto magnetic beads was incubated with purified ssDNA from the positive selection in a 100 μ L volume. After an incubation period, the supernatant was removed and beads were washed three times with 25, 25, and 50 μ L SB, with removed supernatant and washes serving as a PCR template. Amplified dsDNA was then prepared for another selection round as outlined above. Negative selections done in this manner occurred in Rounds 1, 3, 5, and 11.

Round 6 of selection was done as stated above, however the selection buffer consisted of 10% methanol in 1X selection buffer (methanol selection buffer; MSB) to ensure ssDNA binding in ensuing selection rounds. For Rounds 7-11 of positive selection, the purified ssDNA library was incubated with immobilized target as outlined above. However, after three washes, 100 μ L of 1 mM atrazine in MSB was incubated with the beads for decreasing time periods. The ssDNA bound to the immobilized target was separated from the ssDNA in the solution phase and served as a template for PCR amplification. Preparation of the library for the ensuing round of selection was done as noted above.

The Round 8 negative selection was done as described above, however ssDNA was competitively eluted with the pesticides propanil, 2,4-D acid, and malathion (Figure 2.2 e-g). The ssDNA bound to the immobilized target was separated from solution and resuspended, serving as the template for PCR amplification. Rounds 9 and 10 of negative selection were done in the same manner, with DACT and simazine (Figure 2.2 c-d) serving as targets for competitive elution, respectively.

The final round of positive selection, Round 12, was done similar to the competitive elutions above. However, bound ssDNA was eluted with 100 μ L of 100 μ M atrazine in MSB. This eluted ssDNA served as a template for PCR amplification.

2.2.2 Sequencing of Atrazine-Specific MREs

Every third round of selection, the ssDNA library was analyzed for consensus binding sequences by DNA sequencing. The PCR-amplified (using non-biotinylated primers) library was ligated into the pCRII vector (Invitrogen, Carlsbad, CA) and cloned into competent bacteria per manufacturer's instructions. The plasmid was then extracted using the AxyPrep Plasmid Prep Kit (Axygen, Union City, CA) and sent for sequencing (Eurofins MWG Operon, Huntsville, AL) using the M13R primer complementary to a region upstream of the PCR insert in the pCRII vector. Analyses were done on 30-50 random sequences for each round to identify consensus sequence binding families.

2.2.3 Atrazine MRE Binding Assays

From the Round 12 sequences, three sequences were chosen for further characterization. The predicted tertiary structures were obtained using the Mfold DNA web server at 25°C with buffer salt conditions (SantaLucia 1998; Peyret 2000; Zuker 2003). R12.23, R12.28, and R12.57

were chosen and synthesized by Eurofins MWG Operon with an AlexaFluor 488 label for use in binding studies.

Concentrations of 0, 0.25, 0.5, 0.75, 1, 5, 10, 15, 20, and 25 nM MRE were used in saturation binding studies. In a 200 μ L total volume of MSB, 10 μ L of immobilized target was incubated with each concentration for five minutes. Unbound MRE was removed and each incubation was washed five times with 200 μ L MSB, then the immobilized target was resuspended in 200 μ L MSB and heated to 95°C for 10 minutes to denature and elute bound MRE. Eluted ssDNA in MSB was placed in a 96-well microplate and measured in a Synergy 2 microplate reader equipped with a tungsten halogen lamp with excitation wavelength of 485 nm and an emission filter at 520 nm using Gen5 1.06 software (Biotek US; Winooski, VT). All fluorescence readings on the plate were normalized to 100 μ L of a 1 nM solution in SB of the fluorescent MRE and the no ssDNA incubation. For each concentration set, the same was done with washed streptavidin-coated magnetic beads to ensure binding over background to the beads. Each set of incubations were performed in triplicate. To determine the dissociation constant of R12.23, data were analyzed with Origin 8 (OriginLab Corporation; Northampton, MA) using nonlinear regression analysis and fit with the equation, $Y = ((B_{\max} * X) / (K_d + X)) + NS * X$, where B_{\max} is the maximum binding, K_d is the dissociation constant, and NS is nonspecific binding as previously described (Langan et al. 2012).

To determine binding of the selected MRE, R12.23, to negative targets used in the selection, 5 nM fluorescent ssDNA was used. For each elutant used, in a 200 μ L total volume of MSB, 10 μ L of immobilized target was incubated with R12.23 for five minutes. The magnetic beads were washed five times with 200 μ L MSB. Then, each incubation was resuspended in 200

μL of 100 μM of the following in MSB: atrazine, desethyl atrazine, DACT, simazine, propanil, 2-4, D Acid, malathion, and MSB only. An incubation with no R12.23 present and

Table 2.1. Decoy-SELEX scheme for atrazine MRE selection.

Round	Positive Selection	Negative Selection
1	Immobilized Target (IT) 25 hrs	Immobilization Substrate (IS) 23 hrs
2	IT 23 hrs	-
3	IT 18 hrs	IS 18 hrs
4	IT 12 hrs	-
5	IT 10 hrs	IS 16 hrs
6	IT w/ methanol buffer 12 hrs	-
7	IT 10 hrs, Competitive Elution w/ 1 mM Atrazine (CE) 5 min	-
8	IT 2 hrs, CE 2 min	IT 2 hrs, CE w/ propanil, 2,4-D acid malathion, 5 min
9	IT 5 min, CE immediate	IT 5 min, CE w/ DACT 5 min
10	IT 1 min, CE immediate	IT 5 min, CE w/ Simazine 5 min
11	IT immediate, CE immediate	IS 18 hrs
12	IT immediate, CE w/ 100 μM atrazine	-

In vitro selection process for obtaining atrazine-specific MRE. Immobilization target (IT) is desethyl atrazine bound to magnet beads. Immobilization substrate (IS) is streptavidin-coated magnetic beads plus blocked biotin reagent. Competitive elution (CE) is removal of bound ssDNA from target-coated magnetic beads by free pesticide in solution. Times listed are incubation times in hours (hrs) or minutes (min).

with atrazine as the elutant served as normalization controls. Each elutant was incubated with the immobilized target for five minutes, and the solution was removed from the beads. This solution was placed in a 96-well plate and measured in a fluorescence plate reader. Data were normalized to an internal fluorescent standard and then the no R12.23 control was subtracted. Each set of specific binding studies were performed in triplicate. Data were averaged and

standard deviations calculated. For each elutant, a t-test was performed to determine statistical differences in the means. To determine binding to other molecules not used in the selection, the same protocol was used. Following incubation of 5 nM fluorescent R12.23 with immobilized target, each incubation was eluted with 100 uM in MSB of the following: atrazine, propazine, terbuthylazine, sebuthylazine, and bovine serum albumin (BSA), which was used as a model of binding to large, globular proteins.

2.3. Results and Discussion

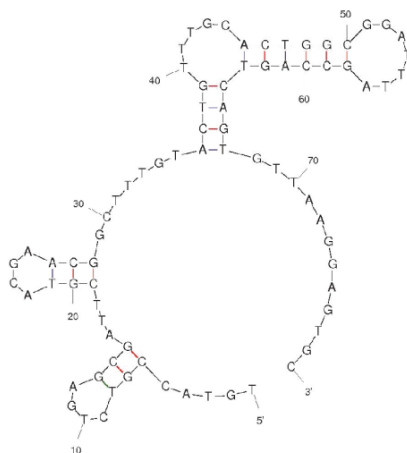
2.3.1 Selection of Atrazine-Specific MREs

Twelve rounds of Decoy-SELEX were completed to obtain ssDNA MREs which specifically bind to atrazine (Table 2.1). The selection scheme ensured that ssDNA MREs will bind to atrazine in solution and neither the immobilization substrate nor malathion, propanil, 2,4-D acid, DACT, or simazine. PCR cycling numbers and ssDNA recovered after preparation and used in the ensuing selection round varied with each round of selection. After every third round of selection (rounds 3, 6, 9, 12 plus after round 10), 30-50 random sequences of the ssDNA molecules in the library were obtained for consensus sequence family analysis. Additionally, random sampling of ssDNA sequences binding to both DACT and simazine was performed. Consensus sequence families were analyzed and tracked throughout the Decoy-SELEX process. From the round 12 library, three sequences were chosen based on their inclusion in consensus sequence families and predicted tertiary structures. Those sequences, R12.23, R12.28, and R12.57 were assayed for their binding affinity to atrazine (Figure 2.3). Those sequences had Gibb's free energy values of -11.32 kcal/mol, -10.71 kcal/mol, and -10.61 kcal/mol, respectively.

A

R12.23	TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
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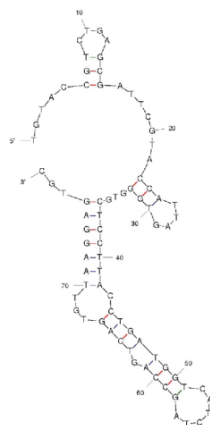
R12.23 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.24 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.34 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC



B

R12.28	TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
--------	---

R12.26 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.28 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.51 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC



C

R12.57	TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
--------	---

R12.28 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.57 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.59 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC
R12.24 TGTACCGTCTGAGCGATTCTGACGAAACGGCTTTGTACTGTTTGCACCTGCGGATTAGCCAGTCAGTGTAAAGGAGTGC

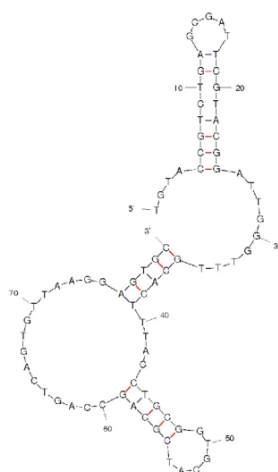


Figure 2.3. Sequences and structures of atrazine MREs. The ssDNA sequence of selected atrazine MREs, which is above a representative consensus sequence family, with the secondary structure as predicted by Mfold below (SantaLucia 1998; Peyret 2000; Zuker 2003). The sequence and structure of (a) R12.23, (b) R12.28, and (c) R12.57.

2.3.2 Affinity and Specificity of Atrazine-Specific MRE

Fluorescent saturation binding assays were performed to determine the affinity of R12.23, R12.28, and R12.57 for atrazine. Concentrations of MRE in the high pM to low nM range were assayed, plotted, and fit by nonlinear regression analysis (Figure 2.4). The equilibrium dissociation constant (K_d) of R12.23 was determined to be between 0.14 nM and 0.97 nM (Table 2.2). This is in the lower range of MREs targeting small molecules and comparable to MREs targeting proteins (Hamula et al. 2006; Stoltenburg et al. 2007). Such high affinity for atrazine is likely due to the multiple rounds of selection with very stringent binding conditions utilizing the Decoy-SELEX process. Fluorescent binding assays were also performed for R12.28, which showed a K_d between 1.28 nM and 9.20 nM, and R12.57, which showed a K_d between 4.42 nM and 51.74 nM. Therefore, because of its high affinity and reproducible dissociation constant range, R12.23 was chosen for further binding analysis.

Further sequence analysis on R12.23 determined possible binding sequences. This sequence had the greatest stability in the round 12 library as predicted by Mfold. It was present six times in four out of eleven total large round 12 sequence families and was not evident in any previous round of selection. Interestingly however, one five-base consensus sequence in R12.23 was also found in a very large round 10 consensus sequence family (GCTTT). Another five-base consensus sequence in R12.23 was found in large families in both round 6 and 9 (TGTTT). This particular five-base sequence was also found as part of a consensus sequence family for ssDNA that bound to both DACT and simazine. A second five-base consensus sequence in R12.23 also had these characteristics (GCACT). These two sequences appeared consecutively in R12.23, and

were part of a large dual-hairpin in the predicted structure (Figure 2.3a). This therefore suggests that while this full MRE sequence did not appear in any prior rounds' random sequencing efforts, components therein were important in binding to atrazine and surviving the Decoy-SELEX process while others were important in conferring specificity.

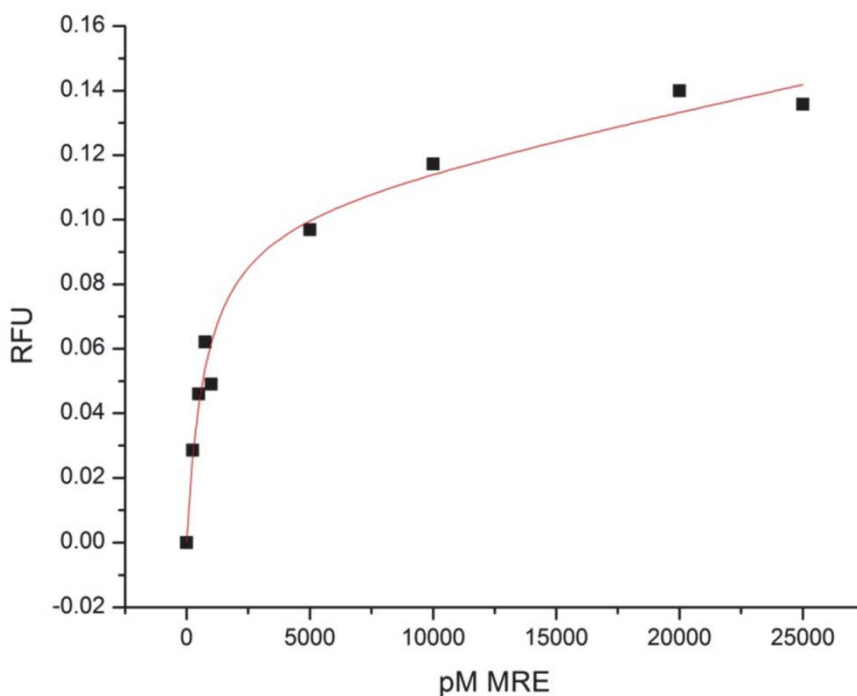


Figure 2.4. Fluorescent equilibrium binding assay of R12.23. Representative saturation binding assay with nonlinear regression best fit. K_d R12.23 for atrazine in this data is 0.75 nM \pm 0.22 nM.

Table 2.2. Equilibrium dissociation constant (K_d) of atrazine MREs.

Plot	R12.23	R12.28	R12.57	R12.23.TRNC1
1	0.75 \pm 0.22 nM	1.28 \pm 0.85 nM	51.74 \pm 7.75 nM	0.079 \pm 0.065 nM
2	0.14 \pm 0.04 nM	9.20 \pm 3.02 nM	6.18 \pm 8.67 nM	0.700 \pm 0.777 nM
3	0.97 \pm 0.61 nM	4.81 \pm 1.75 nM	4.42 \pm 3.67 nM	0.848 \pm 0.826 nM

*All data are presented as $K_d \pm$ Standard Error as given by the binding equation.

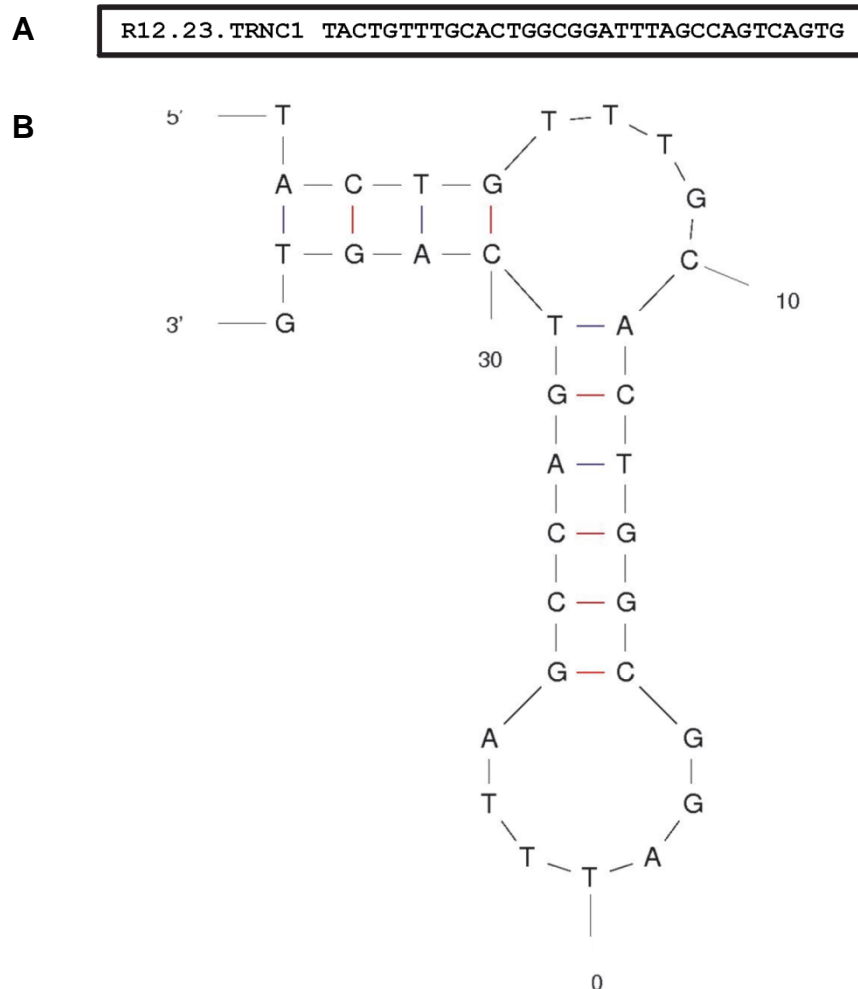


Figure 2.5. Sequence and structure of R12.23.TRNC1. (a) Designed sequence of R12.23.TRNC1. (b) Secondary structure of R12.23.TRNC1 as predicted by Mfold.

To further determine the binding site of R12.23 and in an effort to increase binding affinity, a truncated sequence was designed and termed R12.23.TRNC1 (Figure 2.5). It is comprised of a 34 base sequence which composes the major dual hairpin-loop structural component of the full-length R12.23 discussed in the previous paragraph (Figure 2.5) (synthesized by Katherine Hickey and Peter Gannett of WVU Basic Pharmaceutical Sciences). Fluorescent saturation binding assays using a fluorescein-labeled MRE were performed with this sequence as described above. These assays showed R12.23.TRNC1 had a K_d 0.079 and 0.848

nM (Table 2.2) (data generated by Eamonn Maher and Ginger He of the Sooter Lab). The affinity of the truncated sequence is overall as good as or better than the full-length sequence. This is similar to previous work that has assayed binding affinity of truncated MRE sequences (Kaur and Yung 2012). Therefore, it is likely that this is the binding domain of R12.23.

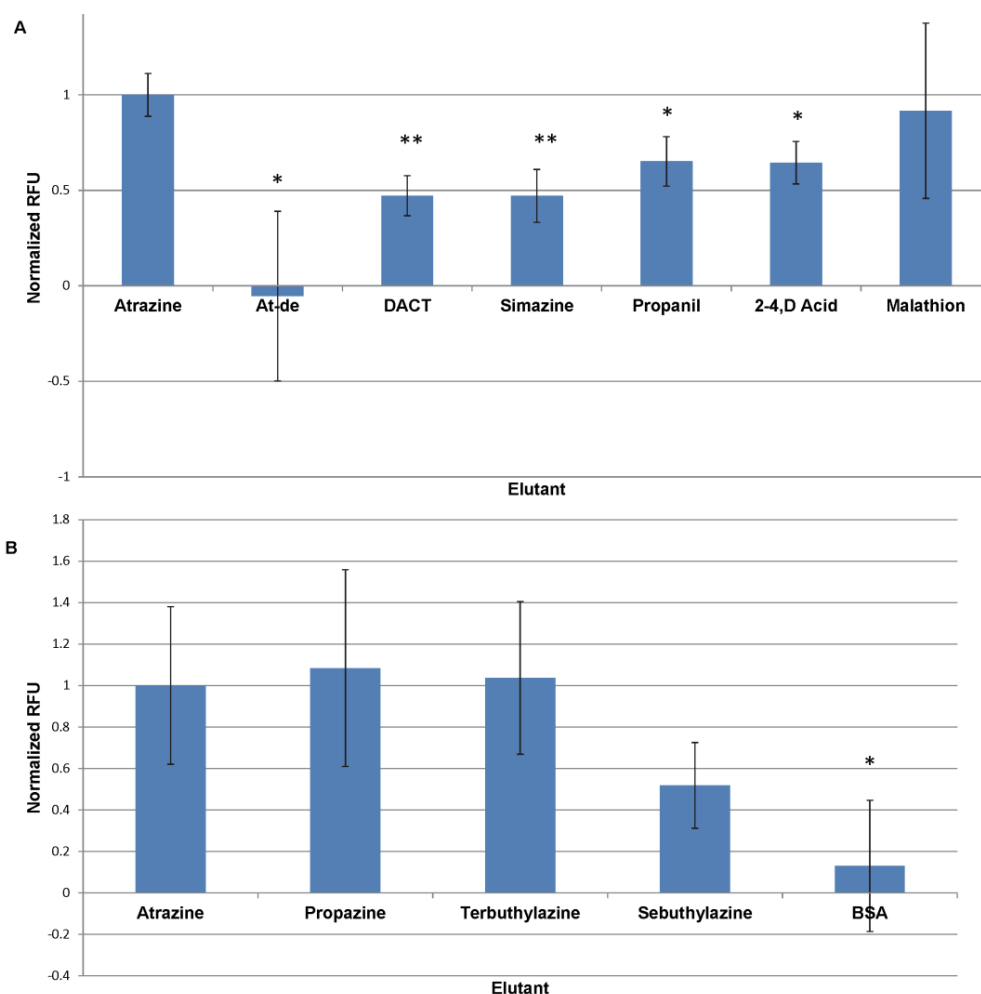


Figure 2.6. Cross-binding studies of R12.23 atrazine MRE to other molecules. Data were normalized to atrazine binding which was set to 100%, with error bars representing standard deviations. Statistical significance levels of $p < 0.05$ are designated by ‘*’, while levels of $p < 0.01$ are designated by ‘**’ (a) Atrazine is the target of selection, desethyl atrazine is the derivative used for immobilization to magnetic beads. DACT and simazine are metabolites of atrazine. Propanil, 2-4, D acid, and malathion are pesticides found in close environmental proximity to atrazine. (b) Propazine, terbutylazine, and sebuthylazine have closely-related chemical structures, but were not used in the selection scheme. Bovine serum albumin (BSA) readily binds many molecules and was used as a model of the MRE binding to proteins.

Fluorescent specificity binding assays were also performed on the atrazine MRE using elution from immobilized target beads. Analysis showed significantly higher binding of R12.23 to atrazine in solution than to four of the five negative targets used in the selection (Figure 2.6a). These included DACT, simazine, propanil, and 2-4,D Acid (Figure 2.2). Binding to malathion in solution did occur but was highly variable. This may be because malathion is known to be reactive toward DNA by cleaving the strand (Pluth et al. 1996; Reus et al. 2008; Moore et al. 2010). This reactivity of malathion may not affect signal of a field- deployable sensor if it is designed to function based on MRE folding and not cleavage. It is important to note that R12.23 is selective for atrazine over simazine, as current molecular recognition elements for atrazine are not (Wortberg et al. 1996). Also, the MRE has no affinity for desethyl atrazine in solution. This is noteworthy because this atrazine derivative was immobilized on magnetic beads and used for the initial positive rounds of selection. Additionally, even though binding to atrazine is approximately two-fold higher than it is to DACT or simazine, the dissociation constant may be much higher than twice that of R12.23 for atrazine. Therefore, incorporation into a sensor may lead to much greater specificity than it appears here. It is clear that stringent competitive elutions selected for an MRE that has high specificity for atrazine in solution compared to the negative targets.

Interestingly, structurally similar molecules (Figure 2.2) which were not used as negative targets bound to the selected MRE (Figure 2.6b). This may be due to the structural complexity of the MRE. Binding is statistically as strong to propazine, terbuthylazine, and sebuthylazine. This suggests that R12.23 requires the chlorotriazine base, as do antibodies that bind to atrazine (Herranz et al. 2008). It also suggests that it requires a certain level of complexity on the amine-linked side-groups of the chlorotriazine base. Based on the structures, it seems binding requires

at least four total carbons in the side-groups. As simazine also meets this requirement, it seems a second requirement is having at least one side-group with three carbons. This is evidenced by the lack of binding to simazine, DACT, and desethyl atrazine, which do not meet these complexity requirements. Propazine, with two amine-linked isopropyl groups, terbuthylazine, with an amine-linked ethyl and an amine-linked tert-butyl, and sebuthylazine, with an amine-linked sec-butyl, exceed these complexity requirements. It is unclear how more complex chlorotriazines would be bound by the MRE. Exceedingly-complex chlorotriazines, however, likely would be too large to fit in the MRE binding pocket. Bovine serum albumin (BSA) was used as an example of potential MRE binding to proteins. The MRE did not show significant binding to BSA. It is likely that, in spite of some cross-reactivity to more complex chlorotriazines, R12.23 will be useful in a sensor for a class of molecules. After rapid, field-based detection with the MRE, more rigorous analytical approaches such as gas chromatography can be used to identify all present contaminants.

The high affinity and specificity of the atrazine MRE are both important characteristics for its further use in detection applications, such as contaminated ecosystems or individuals. Nucleic acid MREs have previously been developed and used in environmental biosensing for small molecules (Strehlitz et al. 2012; Yildirim et al. 2012). They have also been used for detection of analytes in serum (Soh et al. 2009). In both contexts, it is necessary for the MRE to differentiate between the target and closely-related molecules or other contaminants. The novel selection method reported herein is capable of generating such MREs. Once atrazine-polluted environments or contaminated at-risk individuals are identified, they can be actively remediated or treated with chemical, physical or biological methods (Mandelbaum et al. 1995; Saltmiras and Lemley 2002; Herzberg et al. 2004).

2.4 Conclusions

A novel variation of the SELEX process, called Decoy-SELEX, is presented to select a molecular recognition element with high affinity and specificity for the herbicide atrazine. The MRE binds with a sub-nanomolar equilibrium dissociation constant and is selective for atrazine over several metabolites and other pesticides. The results validate the Decoy-SELEX process while identifying a new mechanism for atrazine detection in environmental and biological mediums. This work will improve upon current analysis of atrazine contamination and allow for informed remediation and treatment efforts.

CHAPTER 3

Introduction to the Properties and Applications of Single-Wall Carbon Nanotubes

3.1 Introduction

Carbon nanotubes are cylindrical fullerenes that can be visualized as rolled graphene sheets with walls of single atom thickness. Their discovery stemmed from the discovery of buckminsterfullerene in 1985 (Kroto et al. 1985). Sumio Iijima originally described multi-wall carbon nanotubes (MWCNT) in 1991 (Iijima 1991; Ajayan 1999). Single-wall carbon nanotubes (SWCNT) were independently described by Iijima and Ichihashi as well as an IBM team in 1993 (Bethune et al. 1993; Iijima and Ichihashi 1993). Since this time, research into their production, properties, and applications has increased.

In those original investigations by Iijima the carbon nanotubes were produced by the arc-discharge method. MWCNT were discovered as a deposit on the graphite cathode tip of this chamber (Iijima 1991; Ando and Zhao 2006). In contrast, SWCNT were discovered as a soot from the gas phase of an arc-discharge chamber using either an iron or cobalt catalyst (Bethune et al. 1993; Iijima and Ichihashi 1993; Ando and Zhao 2006). Characterization and confirmation of the products was performed using transmission electron microscopy and electron diffraction. There have since been new production and characterization methods but these remain useful in the field.

Physical differences between MWCNT and SWCNT confer unique properties. The diameter of an individual SWCNT is typically in a narrow range between 0.5 and 2 nm, while that of a MWCNT is 2-25 nm (Ajayan et al. 1999). This is because SWCNT have walls only one atom thick, while MWCNT have concentric walls approximately 0.34 nm apart (Ajayan 1999). Both, however, have a high aspect ratio, with individual tubes typically having lengths in the micron range. Due to variations in how the edges of graphene sheet meet, walls of both carbon nanotube forms may have helicity, which plays a large role in their physical properties. The

walls of MWCNT may have different helicity, and therefore less uniform with less ideal optical and electronic properties for individual tubes (Ajayan 1999). Individual SWCNT species, however, have uniform and characteristic properties which give them greater potential for application. The helicity of a SWCNT species is also referred to as its chirality, or its (n,m) species (Figure 3.1). This designation derives from the angle at which the tube is rolled. Achiral carbon nanotubes are either in the armchair (n,n) or zigzag (n,0) conformations. Some properties have been investigated, however there is much yet to be determined.

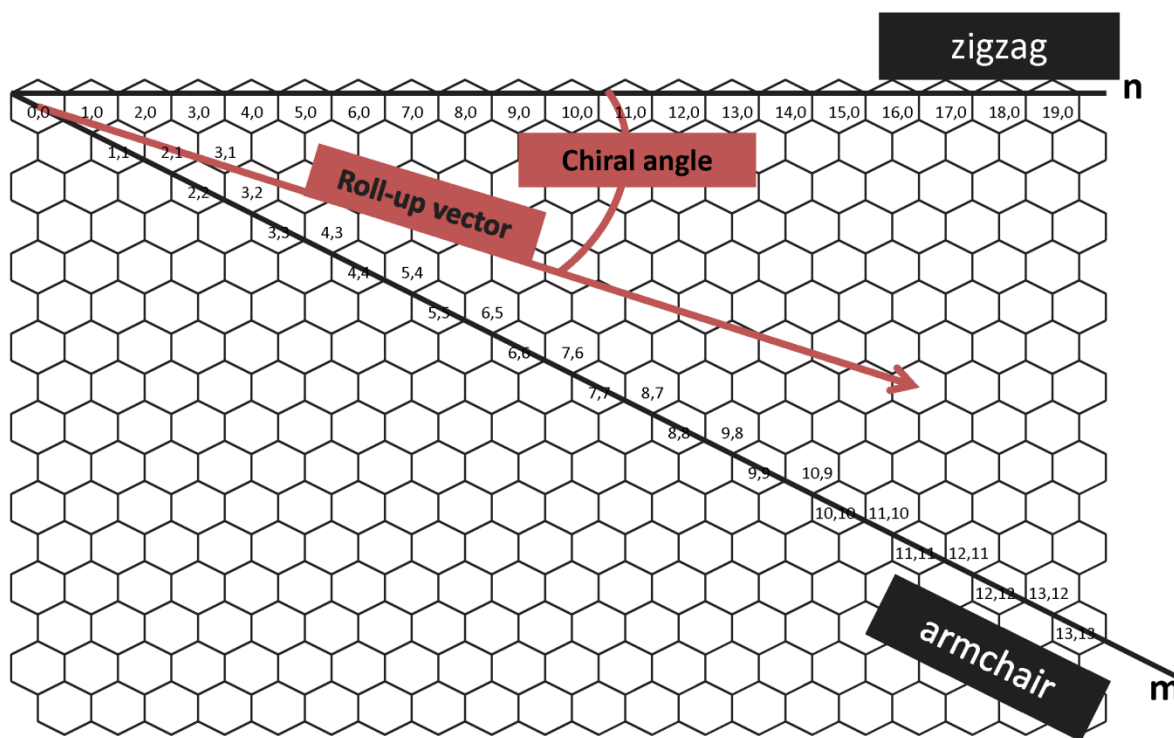


Figure 3.1. Derivation of SWCNT (n,m) chiral species. A visual representation of a graphene sheet depicts the derivation of SWCNT chirality. Carbon bonds form hexagons, and SWCNT can be visualized as rolled-up graphene sheets. The angle at which the sheet is rolled determines the chirality, or the (n,m) designation of the tube. Some n,m species are shown here, with the lines from which zigzag and armchair chiral species are derived. Image is derived from previously-published depiction (Weisman 2010).

The potential application of SWCNT has led to an increase in their production and use. There has been a push to determine their interaction with biological materials. This is because

they have enormous potential in biological sensors and medical treatments. It is also true, however, because there will be an increase in environmental contamination by these particles. It is therefore necessary to understand the production of SWCNT, their special properties, and the applications of SWCNT.

3.2 SWCNT Production

The initial synthesis of SWCNT by the arc-discharge method with a catalyst is still one of the three major forms of their industrial synthesis. Synthesis by laser-ablation and chemical vapor deposition for catalytic growth has since been adopted.

3.2.1 Arc-Discharge Method

The arc-discharge method of SWCNT product was the first developed and remains to be in use because it is simple and has a relatively high yield (Baddour and Briens 2005). To produce SWCNT, a chamber is filled with an inert gas and contains two graphite electrodes (Bethune et al. 1993; Iijima and Ichihashi 1993). A metallic catalyst such as iron or cobalt is placed on either of the electrodes. Then a direct current is applied between the electrodes and SWCNT are deposited from the vapor phase of the chamber. The resulting SWCNT products are relatively short, bundled, and need purification due to impurities (Saifuddin et al. 2013). This method was scaled-up from initial production in 1997 by the work of Journet *et al.* (Journet et al. 1997), however larger-scale production remains difficult.

3.2.2 Laser-Ablation Method

Laser-ablation synthesis of SWCNT is the most expensive method of SWCNT production, however the product is of relatively high purity (Baddour and Briens 2005). A target of graphite with metal catalysts is inserted into a furnace with an inert gas in this method (Guo et al. 1995; Thess et al. 1996). High-intensity laser pulses are then directed at the graphite and

SWCNT are deposited on a water-cooled collector (Popov 2004). This method is only useful for production of SWCNT and not MWCNT. Additionally, it can be more finely tuned by laser and gas properties to produce SWCNT with specific characteristics (Saifuddin et al. 2013).

However, the product is tangled and needs purification. This coupled with the high cost makes this method is not as extensively used as the other two.

3.2.3 Chemical Vapor Deposition Method

Synthesis of SWCNT by chemical vapor deposition (CVD) is the most inexpensive and easiest method to scale up for industrial purposes. This method involves the production of SWCNT by flowing hydrocarbons and an inert gas over a metal catalyst in a furnace (Popov 2004). It was initially used for production of carbon filaments before the discovery of SWCNT (Baker 1989). Since then, however, it has been adapted and scaled up for industrial carbon nanotube production (Joseyacaman et al. 1993; Amelinckx et al. 1994; Ivanov et al. 1994; Li et al. 1996). This process generally favors the production of MWCNTs, however the characteristics of SWCNT are highly controllable under specific conditions (Kong et al. 1998; Baddour and Briens 2005; Saifuddin et al. 2013). Overall, the CVD method typically has high purity and produces longer nanotubes. The only drawback to this method is that the SWCNT typically have many defects (Saifuddin et al. 2013). Despite this, the simplicity, cost-effectiveness, and scalability of the CVD method make it the most widely-used for industrial purposes.

3.2.4 Purification and Enrichment

After synthesis of SWCNT, they may undergo multiple processes before their study and application. These include purification of the nanotubes from synthesis contaminants and

enrichment of particular SWCNT types. Both of these steps may affect the SWCNT properties and neither of them is completely optimized.

Purification of SWCNT is necessary after the synthetic process due to carbonaceous and catalytic metal impurities. The most commonly-used methods for this are oxidation, filtration and chromatography, chemical functionalization, and microwave heating (Haddon et al. 2004; Park et al. 2006). There are, however, other techniques that may be used in conjunction with these, such as centrifugation, ultrasonication, and magnetic separation (Saifuddin et al. 2013).

Another possible form of post-synthetic processing is enriching SWCNT of a particular diameter, length, or chirality. It is important to be able to do this because these characteristics are strong determinants of SWCNT physical properties (Zhang and Zheng 2010). This may be partially done through controlled synthesis with specific temperatures, catalysts, or substrates (Bandow et al. 1998; Kukovitsky et al. 2002). However, it is also possible to enrich particular SWCNT types by oxidation and etching (Zhang et al. 2006; Miyata et al. 2007). Enrichment of SWCNT by chirality and diameter is possible through ultracentrifugation (Arnold et al. 2006). Finally, the affinity of particular surfactants for individual (n,m) species has been used to separate SWCNT (Ju et al. 2008; Marquis et al. 2008). Of these, DNA is the most promising, as individual sequences have greater affinity for particular diameters and chiralities (Huang et al. 2005; Albertorio et al. 2009; Tu et al. 2009; Tu et al. 2011). Obtaining pure SWCNT chiral species is important for many applications, however the process remains difficult to obtain highly pure samples (Komatsu and Wang 2010; Zhang and Zheng 2010).

3.3 Properties of SWCNT

Single-wall carbon nanotubes have many properties that make them ideal for a range of applications (Table 3.1). These include special mechanical, electrical, and optical characteristics.

As mentioned previously, the properties depend strongly on the SWCNT diameter, chirality, and length.

Table 3.1. Special properties of single-wall carbon nanotubes (SWCNT) and their correlation to applications.

Property	Description	Application
Mechanical	<ul style="list-style-type: none"> -Very strong -High stiffness -Flexible -Large surface area -Low density 	<ul style="list-style-type: none"> -Replacement of carbon fibers for material reinforcement -Polymer integration for flexibility and strength
Electrical	<ul style="list-style-type: none"> -Individual chiralities are either metallic or semiconducting 	<ul style="list-style-type: none"> -Conductors in electronics -Signal transducers in sensors
Optical	<ul style="list-style-type: none"> -Fluorescence in the near-infrared region -Exact fluorescence properties dependent on chirality and length 	<ul style="list-style-type: none"> -Optical fluorescence sensors -<i>In vivo</i> imaging -Chirality, length, and dispersion characterization
Functionalization	<ul style="list-style-type: none"> -Ability to be chemically modified -Ability to be wrapped or adsorbed to by biological molecules 	<ul style="list-style-type: none"> -Potential for aiding in all previous applications -Biological sensors

3.3.1 Mechanical Properties

The mechanical properties of SWCNT have made them of high interest for many industrial applications. They are considered very strong with high stiffness due to characteristic sp^2 carbon bonding (Robertson et al. 1992; Popov 2004). Individual SWCNT have an average Young's modulus value of 1.8 TPa, which is an extraordinarily large measure of stiffness for a material (Jacobsen et al. 1995). Important for engineering and industrial purposes, the value for bundled SWCNT is estimated to be 600 GPa; lower due to weaker associations and formations of individual tubes (Salvetat et al. 1999). This, however, can theoretically be up to 38 TPa for identical achiral SWCNT, which could be extremely useful in engineering applications (Popov et

al. 2000). They are unique in that they also have a high flexibility in spite of their strength and stiffness (Coleman et al. 2006). Additionally, SWCNT have a very high surface area and a relatively low density of 0.6 g/cm^3 (Ajayan 1999).

3.3.2 Electrical Properties

Single-wall carbon nanotubes may be metallic or semiconducting based on their chirality. Of all possible SWCNT species, 33% are metallic and 67% are semiconducting (Saito et al. 1992; Zhang and Zheng 2010), with all armchair conformations being metallic and zigzag and chiral being either (Ajayan 1999). These properties make it possible to use individual species or broad classes of SWCNT in electronics or sensors. Additionally, it is possible to use nanotubes to create intramolecular electronic junctions (Ouyang et al. 2002). Overall, the electronic properties of SWCNT have been studied largely by scanning tunneling microscopy (STM) (Ouyang et al. 2002). Due to these properties, SWCNT have a great deal of potential in electronic and sensing applications.

3.3.3 Optical Properties

The specialized optical properties of SWCNT make them particularly important for sensing and biomedical applications. Semiconducting nanotubes have optical absorption peaks corresponding to excitation of electrons from valence to conduction bands, a separation referred to as the band gap (Weisman 2010). These well-defined peaks are referred to as E_{11} , E_{22} , and so on, which fall in the near-infrared (IR) and visible light regions, respectively. Fluorescence has been observed across the near-IR band gap of SWCNT (O'Connell et al. 2002). As this is highly dependent on the diameter and chirality of the SWCNT, it is possible to characterize the nanotubes present in a sample based on its optical properties (Bachilo et al. 2002). Furthermore, the near-IR region is relatively unabsorbed by tissue, thus making SWCNT of great potential for

biological imaging and therapeutics (Welsher et al. 2011). SWCNT can generate and detect light while functioning as a transistor, and therefore have great potential for combined electronic and optical devices (Avouris et al. 2008). It is also possible to use absorbance versus fluorescence as a relative measurement of SWCNT dispersion in solution. This is because bundled SWCNT absorb light, but their fluorescence output is dampened (Weisman 2010). It is therefore obvious that optical characterization and applications of SWCNT have great potential.

3.4 SWCNT Applications

Stemming from these properties of SWCNT is the ability to use them in a variety of applications (Table 3.1). They have been integrated into industrial, electronic, sensing, and biomedical processes, among others. It is also important to note that for many applications, it is necessary to have pure or enriched SWCNT species in order to have uniform properties.

3.4.1 Industrial Applications

Due to their unique mechanical properties, SWCNT have been incorporated into polymers to form composites with desirable properties (Breuer and Sundararaj 2004). They are often used instead of carbon fibers, which are used to reinforce and enhance materials for aerospace, automotive, or sporting and protective equipment. They can also be integrated into composites for their ability to dissipate electrostatic interactions, which can be important in safer materials. Considerable research has gone into optimizing the interface of SWCNT and polymers to extract ideal properties (Thostenson et al. 2001). This work has significantly increased the stiffness of composite materials and led to an increase of SWCNT-related products in the marketplace (Coleman et al. 2006).

3.4.2 Electronic Applications

The application of SWCNT in electronics has also become more widespread. This is because of their excellent conductivity, surface area, and ability to take different macroscopic shapes (Schnorr and Swager 2011). SWCNT have thus been studied for use in lithium ion batteries and transparent electrodes, which are components of organic solar cells and organic light-emitting diodes (OLEDs) (Ng et al. 2005; Kim et al. 2009; Landi et al. 2009; Chien et al. 2010). Additionally, they can be used as capacitors and even to build smaller and more efficient electronic circuits, among many other uses (Franklin et al. 2001; Baughman et al. 2002; Du and Pan 2006; Schnorr and Swager 2011).

3.4.3 SWCNT in Sensors

The use of SWCNT in sensors can be closely related to their electronic applications. However, SWCNT-based sensors may also rely on their optical properties. An example of an electronic-based SWCNT sensor is a field effect transistor (FET) (Tans et al. 1998). The nanotubes can adsorb, be wrapped by, or covalently bound to sensing molecules, which changes the conductance of the nanotube upon binding (Pengfei et al. 2003; So et al. 2008; Schnorr and Swager 2011). They have also been used for detection of gases and chemical vapors through chemically reactive coatings (Novak et al. 2003; Snow et al. 2005). As mentioned above, polymers or other biological sensing molecules can be attached to SWCNT. This includes adsorption or the covalent attachment of antibodies as well as covalent attachment or wrapping of DNA around nanotubes (Zhang et al. 2009). These can function in sensors as described above, or can be part of optical SWCNT sensors (Huang et al. 2011). An important example of this is the near-infrared detection of analytes with DNA and DNA molecular recognition

element(MRE)-wrapped SWCNT (Jeng et al. 2006; Chen et al. 2009). These are among many other examples of research being done to develop SWCNT-based sensors.

3.4.4 Biomedical Applications

One area of great potential for SWCNT use is in biomedical applications, which draws on multiple properties of this material. An important necessity of this use is that the nanotubes are not toxic to humans. Multiple studies have shown that SWCNT functionalized or non-covalently altered with biological molecules reduce their toxic effects (Vardharajula et al. 2012). Therefore, they have been developed for use as *in vitro* and *in vivo* imaging tools, as their near-infrared fluorescence is detectable through cells and tissue (Welsher et al. 2011; Kang et al. 2012). Additionally, they have been used to deliver small molecule and biologic drugs as well as genes to specific diseases (Pantarotto et al. 2004; Portney and Ozkan 2006; Bhirde et al. 2010). Research is ongoing to ensure the safety of SWCNT in biomedical applications, which will lead to an invaluable tool for clinical diagnostics and therapeutics.

3.5 Conclusions

Single-wall carbon nanotubes are an extraordinary and relatively new class of materials into which research is ongoing and increasing constantly. This is due to their unique and desirable properties, which are drawn from their production, purification, and enrichment. They have particular value in both biomolecular sensing and biomedical applications. Both of these uses require further work into understanding the interactions of SWCNT and biological molecules and systems.

The work described here was completed to further determine specific properties of SWCNT. As noted, their interaction with DNA is of particular interest for sorting and separation, sensors, and other applications. The effect of multiple chiralities of DNA wrapped

with different DNA sequences on the polymerase chain reaction was determined. This will allow for future work in separation of SWCNT and in sensors. Additionally, the effect of SWCNT on a green algae model organism was determined. This will help to determine future environmental impacts of widespread DNA-wrapped SWCNT use and whether these constructs are safe for use. Together, these investigations into the interactions of biological systems and DNA-wrapped SWCNT have elucidated properties of SWCNT important for their future use.

CHAPTER 4

Investigation of the Effects of DNA-dispersed Single-Wall Carbon Nanotubes on the Polymerase Chain Reaction

4.1 Introduction

Single-wall carbon nanotubes (SWCNT) are cylindrical fullerenes with walls of single carbon atom thickness (Iijima 1991; Iijima and Ichihashi 1993; Ajayan 1999). They have enormous potential in industrial, sensing, and biomedical applications due to their unique physical, electronic, and optical properties (Heller et al. 2005; Wang 2005; Coleman et al. 2006; Huang et al. 2011). SWCNT may be either metallic or semiconducting, with 33% and 67%, respectively, of all potential species having those characteristics (Saito et al. 1992; Zhang and Zheng 2010). The unique properties of SWCNT are dependent upon the chirality and diameter of the SWCNT, however purification of individual species remains difficult (Zhang and Zheng 2010). Individual types of semiconducting SWCNT have unique optical fluorescence properties (O'Connell et al. 2002). Therefore, in applying SWCNT, it is possible to take advantage of their electrical or optical properties, separately or combined (Avouris et al. 2008).

Single-stranded DNA has been used to disperse hydrophobic SWCNT in solution (Zheng et al. 2003). The interaction of DNA sequences with SWCNT may have implications in sensing and purification techniques. Individual sequences have been identified that preferentially wrap certain SWCNT chiralities (Huang et al. 2005; Albertorio et al. 2009; Tu et al. 2009; Tu et al. 2011). DNA-associated SWCNT have also been used in molecular sensing (So et al. 2005; Jeng et al. 2006; So et al. 2008). The potential use of a DNA-wrapped SWCNT (DNA:SWCNT) platform shows promise in multiple applications; however research is continuing to optimize use of these complexes.

In sensing applications and analysis of environmental samples, it is useful to employ polymerase chain reaction (PCR) techniques in order to exponentially amplify the SWCNT-associated DNA sequence. Additionally, there has been work to develop PCR-based

microfluidic sensing devices (Khandurina et al. 2000), as well as DNA-functionalized SWCNT sensing devices (So et al. 2005). Integration of these two sets of technologies would create new potential analyte detection mechanisms. Therefore, it is important to determine the effect of DNA-associated SWCNT on the polymerase chain reaction.

The aim of this study was to investigate amplification of DNA by the polymerase chain reaction (PCR) in the presence of DNA-dispersed SWCNT. Factors investigated include DNA sequence-dependence, SWCNT chirality-dependence, and DNA:SWCNT concentration-dependence; amplification of the wrapping sequence itself; and age of the DNA-SWCNT preparation. This work will provide a basis for further DNA-dispersed SWCNT studies which require PCR amplification.

4.2 Methods

4.2.1 DNA-Dispersed SWCNT Preparation

DNA:SWCNT preparations were made similarly to previously described methods (Zheng et al. 2003; Lahiji et al. 2007). Briefly, 2 mg CoMoCAT SWCNT were mixed with 2 mg DNA in 2 mL phosphate-buffered saline (PBS). SWCNT used were: without chiral purification “Mix” or (6,5) chirality-enriched (Sigma-Aldrich; St. Louis, MO). DNA sequences used were: estradiol (E2) Molecular Recognition Element (MRE) (Table 4.1) (Kim et al. 2007), TG15 (Table 4.1) (Eurofins MWG Operon; Huntsville, AL), or salmon testes genomic DNA (Sigma-Aldrich; St. Louis, MO). This mixture was sonicated on ice for two hours with a Virsonic XL2020 ultrasonic liquid processor (Misonix; Farmingdale, NY) equipped with a 3.2 mm microtip at approximately 20% power. The resulting solution was centrifuged at 16,000xg for 90 minutes to remove insoluble SWCNT. The solution was then filtered through an Amicon Ultra-4 Centrifugal Filter unit with 100,000 molecular weight filter (Millipore; Billerica, MA) three times at 4000 RPM for 15 minutes each to remove DNA that did not associate with the SWCNTs. The recovered DNA:SWCNT solution was brought to a volume of 500 μ L with PBS. The concentration of the DNA:SWCNT sample was obtained by gravimetric analysis of 10 μ L of the solution

weighed on an Ohaus Discovery Microbalance (Ohaus; Parsippany, NJ). Samples were stored in screw-top containers at room temperature before use. To determine the effect of DNA:SWCNT preparation age on PCR, samples were used that were prepared between four and ten months before assayed or less than one month before assayed. In total, 12 DNA:SWCNT samples were prepared.

Table 4.1. Synthesized DNA sequences used in DNA:SWCNT PCRs.

Sequence Name	DNA Sequence Used
Estradiol (E2) MRE	5'-GCTTCCAGCTTATTGAATTACACGCAGAGGGTAGCGGCTCTGCGCAT TCAATTGCTGCGCGCTGAAGCGCGGAAGC-3'
TG15	5'-TGTGTGTGTGTGTGTGTGTGTGTGTGTGTGTG-3'
F.E2	5'-GCTTCCAGCTTATTGAATTACACGCAGAGGGTAGC-3'
R.E2	5'-GCTTCCGCGCTTCAGCGCGC-3'

4.2.2 SWCNT Characterization

Absorption and near-infrared (NIR) fluorescence measurements were performed as previously described with few modifications (Weisman 2010). Briefly, each DNA:SWCNT preparation was diluted to a 1% solution with distilled-deionized water (Millipore; Billerica, MA). This solution was added to a 10 mm path length cuvette (Starna Cells; Atascadero, CA) and analyzed on a NanoSpectralyzer 1 (NS1) (Applied NanoFluorescence; Houston, TX). The absorbance spectrum was collected from 400 - 1600 nm. Additionally, fluorescent emission spectra were measured using excitation wavelengths of 638 nm, 690 nm, and 784 nm. The resulting spectra were analyzed with ANFSoft (Applied NanoFluorescence; Houston, TX). Fluorescence efficiency measurements, which are fluorescence intensity (RFU) collected across the emission spectrum for a single wavelength divided by the absorbance spectrum, are reported as a relative measure of SWCNT dispersion in solution.

4.2.3 Polymerase Chain Reaction in the Presence of DNA-Dispersed SWCNT

In total, 16 sets of PCR experiments were performed involving 7 individual PCR reactions per set. PCR reactions were composed of: 60 nM E2 MRE, 400 nM forward and reverse E2 MRE primers (F.E2 and R.E2) (Table 4.1), 250 μ M deoxyribonucleotide triphosphates, 1X GoTaq Reaction Buffer (Promega; Madison, WI), 3.5 units *Taq* DNA Polymerase, DNA:SWCNT suspension, and Milli-Q water

to 50 μ L. DNA:SWCNT were added to final concentrations of: 0.01 mg/mL, 0.1 mg/mL, 1 mg/mL, 5 mg/mL, 10 mg/mL. For each set, a negative control was performed with no E2 MRE or DNA:SWCNT and a positive control was performed with no DNA:SWCNT added. Reactions were performed on an MJ Mini Thermal Cycler (Bio-Rad; Hercules, CA) with conditions: initial denaturation of 95°C for 5 minutes; 20 cycles of 95°C for 1 minute 30 seconds, 68°C for 45 seconds, and 72°C for 1 minute 30 seconds; and final extension of 72°C for 7 minutes. Amplification of DNA directly associated with SWCNT was performed using E2 MRE:SWCNT preparation with no additional E2 MRE added to the reaction mix.

4.2.4 PCR Analysis by Agarose Gel Electrophoresis

Each set of control and DNA:SWCNT PCRs was run on a 4% agarose gel for 30 minutes at 135 volts in 1X Tris:Borate:EDTA buffer. Five μ L of each PCR and 3 μ L of 100bp and 25bp ladder (Promega), were mixed with 2 μ L 6X loading dye (Promega) prior to loading in the well. Gels were pre-stained with 5 μ g/mL ethidium bromide and imaged with UV illumination on a BioRad Gel Doc System and analyzed on Quantity One Software (Bio-Rad; Hercules, CA).

4.3 Results and Discussion

4.3.1 Analysis of DNA:SWCNT Preparations

All DNA:SWCNT samples were analyzed by absorbance and NIR fluorescence spectroscopy to determine their SWCNT chirality composition and dispersion in solution (singly-dispersed SWCNTs) (Figures 4.1 and 4.2). DNA:SWCNT samples greater than four months old were more well-dispersed than the identical DNA:SWCNT preparation of less than one month. This would suggest that dispersion of SWCNT in solution with DNA increases with time. Additionally, for both old and new samples, TG15 and E2 MRE dispersed the (6,5)-enriched SWCNT better than mixed chirality SWCNT (Figures 4.1 and 4.2, fluorescence efficiency). These results are consistent with previously published results that have shown short,

synthetic DNA sequences rich in thymine and guanine have a high affinity for (6,5) chiral SWCNTs (Kim et al. 2008). Conversely, the salmon testes genomic DNA:Mix SWCNT samples were more well-dispersed than Salmon testes genomic DNA:(6,5) SWCNT. These results suggest that although the genomic DNA shows the highest association with (6,5)-chiral SWCNTs (Figures 4.1 and 4.2, topography plots) it also forms stable associations with other chiralities.

4.3.2 Agarose Gel Analysis of Old DNA:SWCNT PCRs

Agarose gel analysis of PCRs performed with increasing concentrations of DNA:SWCNT preparations between four and ten months old suggest that accurate amplification products of 76 base pairs (bp) can be obtain at concentrations of 0.1 mg/mL and below for all preparations (Figure 4.1 agarose gels). In most cases, efficiency of PCR amplification is either unaffected or slightly aided at 0.01 and 0.1 mg/mL compared to the positive control reaction (Figure 4.1 relative intensities). The exception to this is the TG15:(6,5) sample (Figure 4.1b), where a very faint band is produced at 0.1 mg/mL. Additionally, bands are produced at 1 mg/mL for TG15:Mix and E2:(6,5) PCRs. No distinct bands are produced at 5 or 10 mg/mL concentration PCRs for any preparation. Prior work has shown that SWCNT sonicated in water just before being added to a PCR reaction increased amplification efficiency of a 410 bp product at concentrations up to 3 mg/mL (Cui et al. 2004). The same study showed that SWCNT in a PCR reaction interacts with both the DNA template as well as *Taq* DNA Polymerase. This study, however, has DNA wrapped around SWCNT, so this is unlikely to be the source of inhibition. Another study showed that that SWCNT functionalized with carboxylic groups produce stronger interactions with *Taq* and more inhibition of PCR than did pristine SWCNT at 0.1-0.8 mg/mL (Yi et al. 2007). Therefore, it is possible there is some interaction between

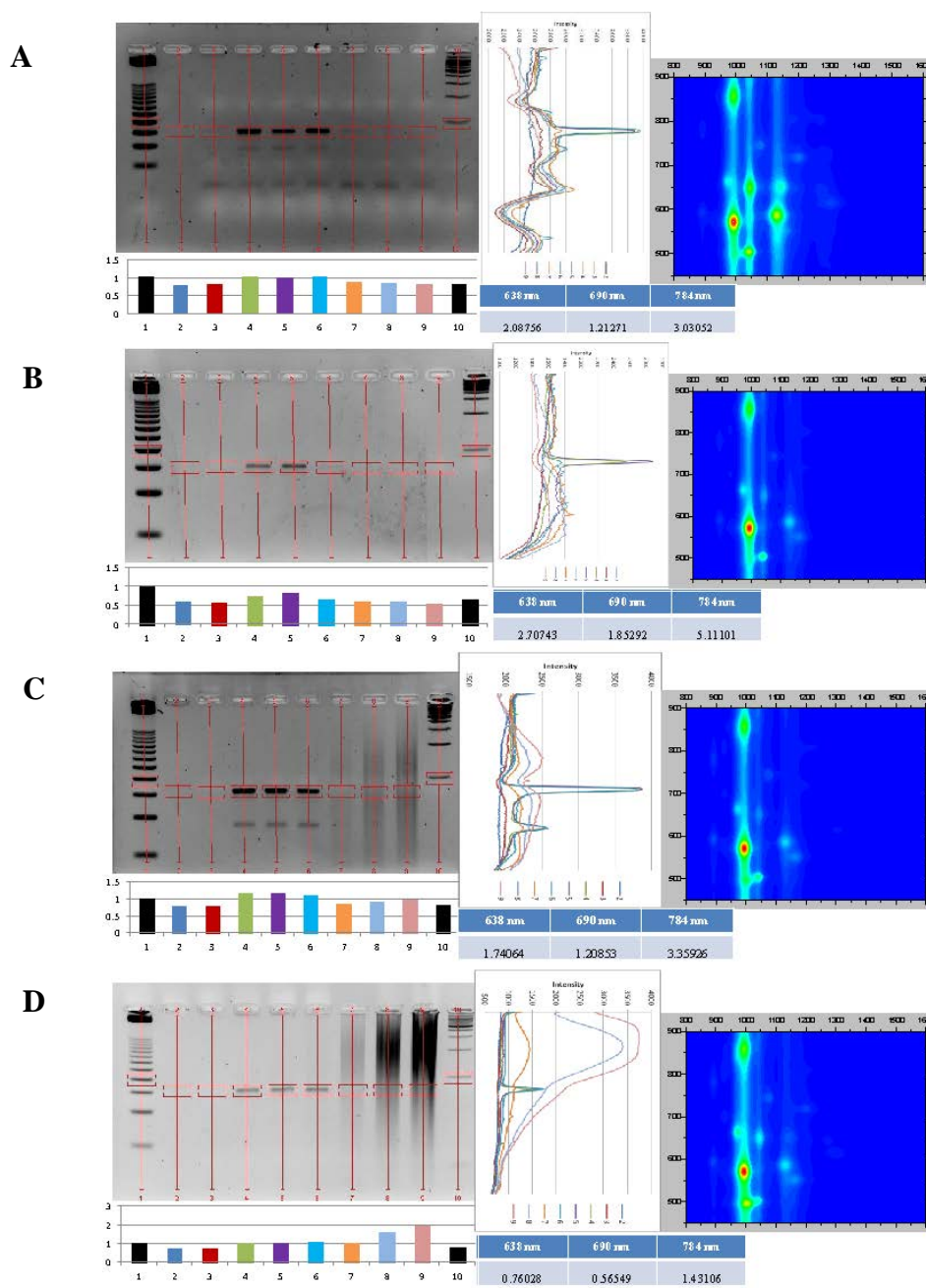


Figure 4.1. (a-d). Agarose gel electrophoresis images and analysis of PCRs with old (4-10 months) DNA:SWCNT samples. For each set, top left is agarose gel image with lanes and quantitized boxes shown, top center is intensity line graph, top right is NS1 synthesized fluorescence intensity plot of DNA:SWCNT samples (y-axis is excitation wavelength, x-axis is emission wavelength), bottom left is bar graph of relative intensity within boxes on gel (normalized to 100 bp marker of 25 bp ladder), bottom center is fluorescence efficiency at each wavelength in NS1 analysis (higher numbers correlate with greater dispersion). Lanes in agarose gel images and relative intensity bar graph: 1. 25 bp ladder (black) 2. empty (dark blue) 3. negative control (red) 4. positive control (green) 5. 0.01 mg/mL DNA:SWCNT (purple) 6. 0.1 mg/mL (neon blue) 7. 1 mg/mL (orange) 8. 5 mg/mL (light blue) 9. 10 mg/mL (pink) 10. 100 bp ladder (black). Intensity line graph colors correlate with bar graph colors in relative intensity graph. (a) TG15:Mix (b) TG15:(6,5) (c) Salmon:Mix (d) Salmon:(6,5).

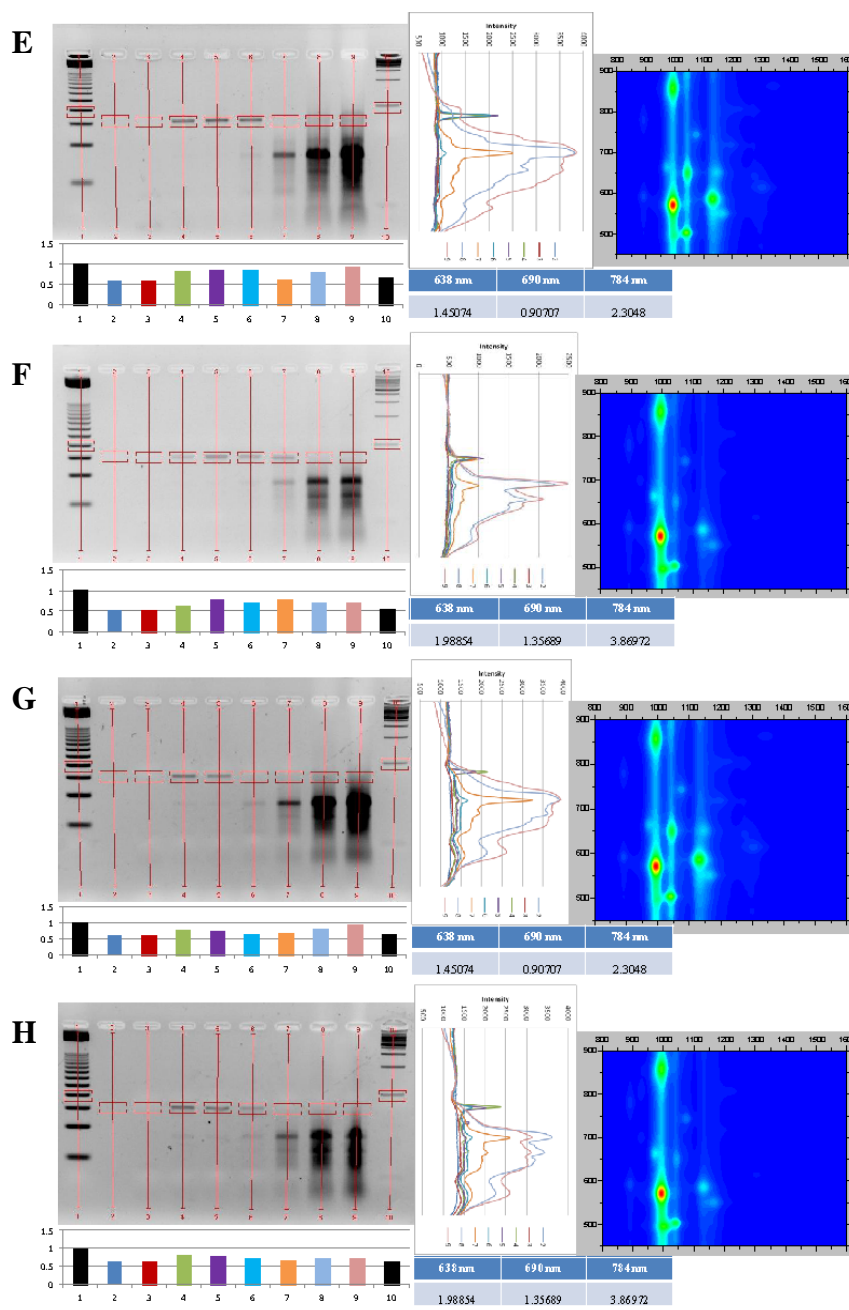


Figure 4.1. (e-h). Agarose gel electrophoresis images and analysis of PCRs with old (4-10 months) DNA:SWCNT samples. For each set, top left is agarose gel image with lanes and quantized boxes shown, top center is intensity line graph, top right is NS1 synthesized fluorescence intensity plot of DNA:SWCNT samples (y-axis is excitation wavelength, x-axis is emission wavelength), bottom left is bar graph of relative intensity within boxes on gel (normalized to 100 bp marker of 25 bp ladder), bottom center is fluorescence efficiency at each wavelength in NS1 analysis (higher numbers correlate with greater dispersion). Lanes in agarose gel images and relative intensity bar graph: 1. 25 bp ladder (black) 2. empty (dark blue) 3. negative control (red) 4. positive control (green) 5. 0.01 mg/mL DNA:SWCNT (purple) 6. 0.1 mg/mL (neon blue) 7. 1 mg/mL (orange) 8. 5 mg/mL (light blue) 9. 10 mg/mL (pink) 10. 100 bp ladder (black). Intensity line graph colors correlate with bar graph colors in relative intensity graph. (e) E2 MRE:Mix (f) E2 MRE:(6,5) (g) E2 MRE:Mix, No MRE (h) E2 MRE:(6,5), No MRE.

DNA:SWCNT and *Taq*, but it is unlikely to be as strong of an interaction and inhibitory effect as with carboxylic SWCNT.

It is also possible to PCR amplify E2 MRE target which is only present in the reaction wrapped around SWCNT (Figure 4.1g and h). These concentration-dependence patterns follow a similar pattern to reactions performed with the addition of free E2 MRE. This is important in that it shows the ability to directly amplify DNA which is associated with SWCNT.

It is also clear that addition of DNA:SWCNT to PCR reactions has effects on amplicons produced. In both Salmon:SWCNT samples (Figure 4.1c and d intensities), smears are visible in lanes which there is no distinct E2 MRE band. This is likely due to degradation of the genomic DNA present in the sample. In both E2:SWCNT samples amplification of a product just less than 50 bp increases with E2:SWCNT concentration. This is also true for E2:SWCNT PCRs performed with no free E2 MRE in the reaction (Figure 4.1e and f). This is likely due to an excess of the amplification target itself in the reaction. However, in the TG15:SWCNT PCRs, no amplicons appeared that were not present in the control reaction.

4.3.3 Agarose Gel Analysis of New DNA:SWCNT PCRs

Analysis of PCRs performed with DNA:SWCNT samples made less than a month before use show amplification in the presence of 0.01 mg/mL (Figure 4.2 agarose gels). At this concentration, all PCRs performed in the presence of free E2 MRE showed an increase in efficiency of amplification (Figure 4.2 relative intensities). However, in contrast to PCRs with old DNA:SWCNT samples, no amplification of E2 MRE was seen at or above concentrations of 0.1 mg/mL. This interesting result suggests that more well-dispersed SWCNT, as seen in the older samples, is less inhibitory to PCR reactions. This may be because more agglomerated SWCNT hinder the reaction ingredients from encountering each other. This is supported by the

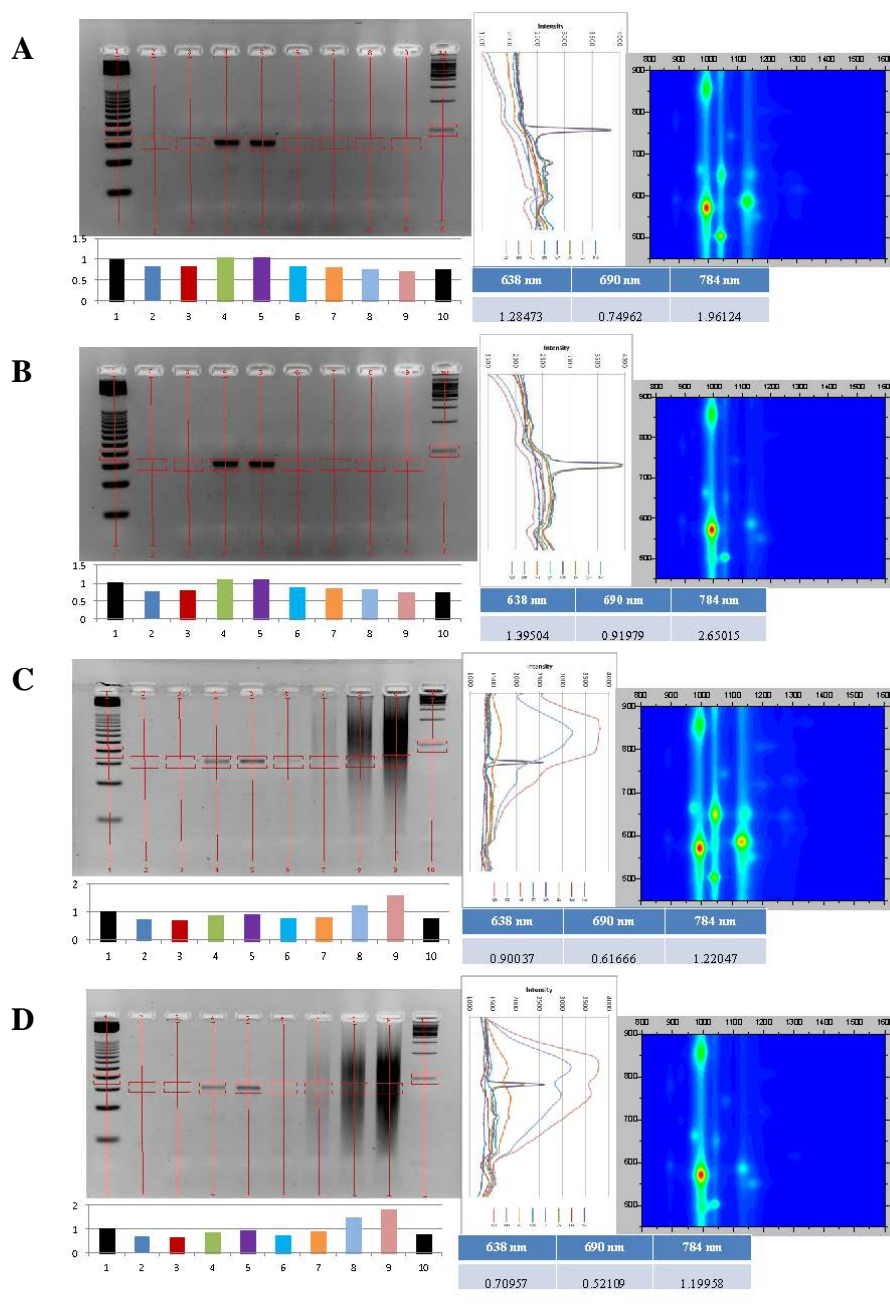


Figure 4.2. (a-d). Agarose gel electrophoresis images and analysis of PCRs with new (<1 month) DNA:SWCNT samples. For each set, top left is agarose gel image with lanes and quantized boxes shown, top center is intensity line graph, top right is NS1 synthesized fluorescence intensity plot of DNA:SWCNT samples (y-axis is excitation wavelength, x-axis is emission wavelength), bottom left is bar graph of relative intensity within boxes on gel (normalized to 100 bp marker of 25 bp ladder), bottom center is fluorescence efficiency at each wavelength in NS1 analysis (higher numbers correlate with greater dispersion). Lanes in agarose gel images and relative intensity bar graph: 1. 25 bp ladder (black) 2. empty (dark blue) 3. negative control (red) 4. positive control (green) 5. 0.01 mg/mL DNA:SWCNT (purple) 6. 0.1 mg/mL (neon blue) 7. 1 mg/mL (orange) 8. 5 mg/mL (light blue) 9. 10 mg/mL (pink) 10. 100 bp ladder (black). Intensity line graph colors correlate with bar graph colors in relative intensity graph. (a) TG15:Mix (b) TG15:(6,5) (c) Salmon:Mix (d) Salmon:(6,5).

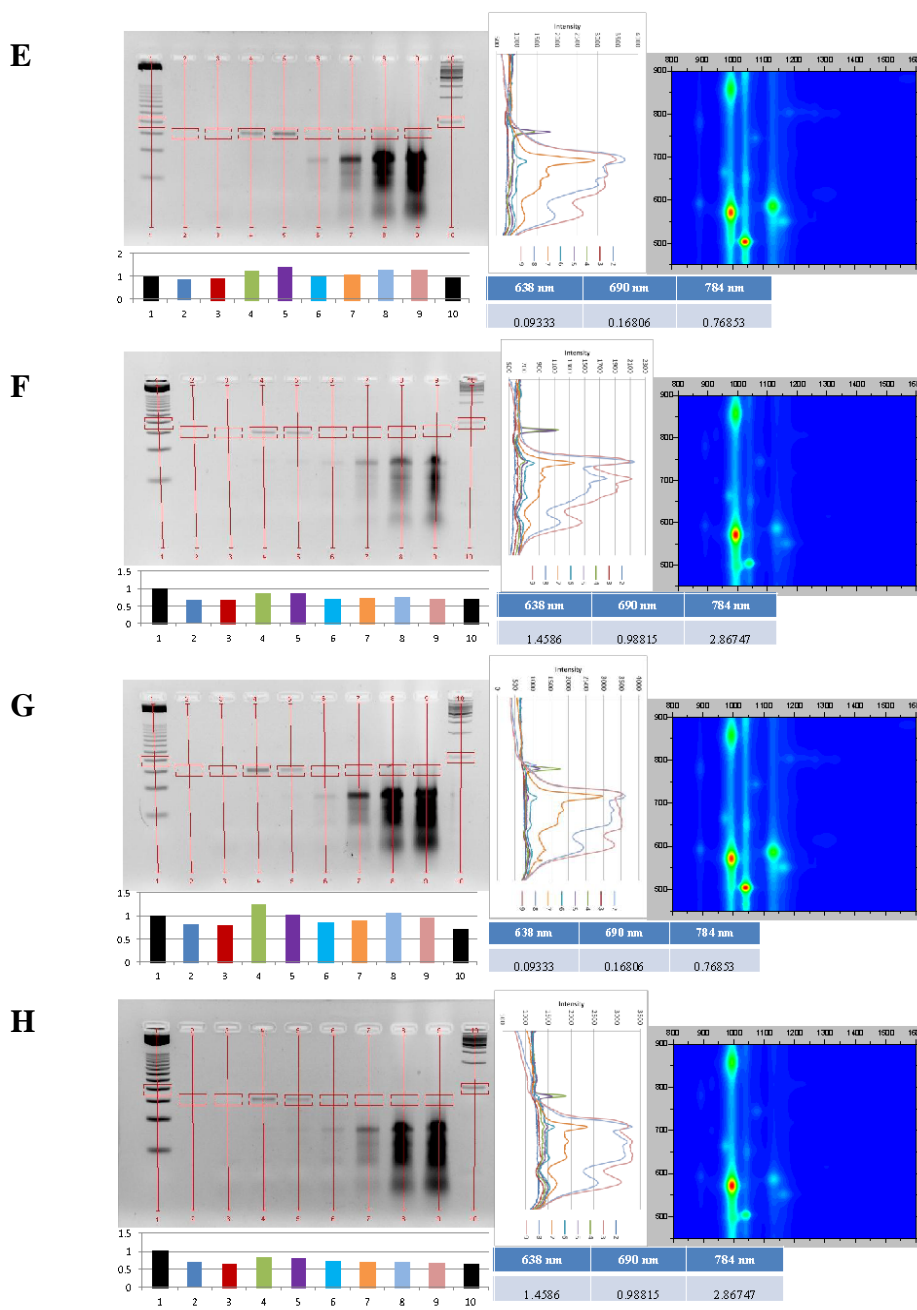


Figure 4.2. (e-h). Agarose gel electrophoresis images and analysis of PCRs with new (<1 month) DNA:SWCNT samples. For each set, top left is agarose gel image with lanes and quantitized boxes shown, top center is intensity line graph, top right is NS1 synthesized fluorescence intensity plot of DNA:SWCNT samples (y-axis is excitation wavelength, x-axis is emission wavelength), bottom left is bar graph of relative intensity within boxes on gel (normalized to 100 bp marker of 25 bp ladder), bottom center is fluorescence efficiency at each wavelength in NS1 analysis (higher numbers correlate with greater dispersion). Lanes in agarose gel images and relative intensity bar graph: 1. 25 bp ladder (black) 2. empty (dark blue) 3. negative control (red) 4. positive control (green) 5. 0.01 mg/mL DNA:SWCNT (purple) 6. 0.1 mg/mL (neon blue) 7. 1 mg/mL (orange) 8. 5 mg/mL (light blue) 9. 10 mg/mL (pink) 10. 100 bp ladder (black). Intensity line graph colors correlate with bar graph colors in relative intensity graph. (e) E2 MRE:Mix (f) E2 MRE:(6,5) (g) E2 MRE:Mix, No MRE (h) E2 MRE:(6,5), No MRE.

fact that higher concentrations have an inhibitory effect on PCR efficiency. However, in both old and new samples, there seems to be no difference in PCR efficiency between Mix or (6,5)-enriched SWCNT samples. This is in spite of a difference in dispersion based on DNA sequence.

In PCR experiments with new E2:SWCNT samples and absent of free E2 MRE, the MRE was amplified from the SWCNT scaffold (Figure 4.2g and h. Amplification of the 76 bp product is observed in the presence of 0.01 mg/mL E2:SWCNT, but not at higher concentration. This follows the same pattern seen as PCRs done in the presence of free E2 MRE. There is, however, a reduction in efficiency at this concentration, which contradicts amplification of free E2 MRE at this concentration.

Amplicons produced from PCRs in the presence of new DNA:SWCNT followed the same general pattern as those with old samples. Previous studies have shown SWCNT increase the specificity of long PCR (14 kb) up to the point that the reaction is inhibited (Zhang et al. 2008). The difference in results may be a function of size of the reaction product, or may be due to the method of SWCNT dispersion, which was sonication in water. Another study has shown that different concentrations of carbon nanopowder (CNP) does not affect amplicons produced in a PCR reaction (Zhang et al. 2007). Therefore, the effects are likely due to degradation of genomic DNA or excess target present in the reaction as previously discussed.

4.4 Conclusions

This work shows differential inhibitory concentrations for old and new DNA:SWCNT preparations of varying dispersion levels in PCR reactions of 0.1 and 0.01 mg/mL, respectively. This work also shows that it is possible to amplify a DNA sequences that is directly associated

with SWCNTS through sonication-mediated dispersion. This is important for purification of SWCNT chiralities using DNA as well as use of DNA:SWCNT-based molecular sensors and environmental sample testing applications.

CHAPTER 5

Investigation of the Effects of DNA-Dispersed Single-Wall Carbon Nanotubes and Sodium Cholate-Dispersed Single-Wall Carbon Nanotubes on the Green Algae *Chlamydomonas reinhardtii*

5.1 Introduction

The increased use of manufactured nanomaterials has led to a need for a better understanding of their potential effects on natural systems. Among these products are carbon nanotubes, which can be represented by sheets of cylindrically-rolled graphene, originally described in 1991 by Iijima (Iijima 1991; Ajayan 1999). The widespread commercialization of single-wall carbon nanotubes (SWCNT) is due to their enormous potential in industrial, biomedical, and electronic sensing applications. This potential is derived from their unique physical, electronic, and optical properties (Wang 2005; Coleman et al. 2006; Huang et al. 2011). The environmental release and transport of SWCNT at all stages of their production and use will lead to environmental concerns (Köhler et al. 2008; Petersen et al. 2011). It is because of their potential that it is necessary to study their interaction with organisms in the environment.

Previous work has shown possible toxic effects of carbon nanomaterials on model organisms. Graphite, C60 fullerene, SWCNT, and multi-wall carbon nanotubes (MWCNT) have been shown to inactivate various Gram-negative and Gram-positive bacteria (Kang et al. 2009). This was shown to increase with incubation times up to four hours. While in the aforementioned work the SWCNT contacted bacteria on a solid support, SWCNT dispersed in solution with various surfactants have shown to be toxic to both bacterial types (Liu et al. 2009). In the model organism *Daphnia magna*, SWCNT suspended in water with no surfactant were the most toxic among other carbon nano- and bulk- materials (Zhu et al. 2009). These previous works make it clear that SWCNT may have toxic environmental effects, but further exploration of various preparations is necessary.

The unicellular green alga *Chlamydomonas reinhardtii* has been developed as a model for the effects of nanomaterials on photosynthetic and microorganisms, as well as an indirect

assessment of ecosystem-level effects (Harris 2001). Work using this model organism has shown toxicity of silver nanoparticles, quantum dots, and titanium dioxide nanoparticles (Navarro et al. 2008; Wang et al. 2008). A subsequent study has shown that this species takes up quantum dots, which cause changes in its transcription profile (Domingos et al. 2011). Additionally, *C. reinhardtii* has been used in finding toxic effects of SWCNT in solution with no surfactant (Matorin et al. 2010). This work showed a negative effect on growth and photosynthetic output of these algae due to SWCNT aggregates in solution. It specifically tested concentrations of 2 $\mu\text{g/mL}$ for SWCNT and 1 and 2 $\mu\text{g/mL}$ of MWCNT, finding varying negative effects of each. Thus, *C. reinhardtii* is an established system for studying the direct toxic effects of nanomaterials and for inferring environmental impact of their release.

This study aimed to determine the effects of well-dispersed SWCNT on the model species *Chlamydomonas reinhardtii*. The system was designed to mimic a stagnant algal growth environment that is similar to natural systems. This design is ideal for well-dispersed SWCNT, however non-dispersed SWCNT would agglomerate and form a precipitate in such a system. Thus, only well-dispersed SWCNT were utilized, mimicking the environmental release of some SWCNT (Köhler et al. 2008). Additionally, agglomerated SWCNT are known to become more well-dispersed over time through interaction with natural organic matter (Lou et al. 2011; Schierz et al. 2012). Therefore, the effects of a wide range of well-dispersed SWCNT in solution with algal growth media on growth and chlorophyll content of green algae was assessed. Furthermore, differences in surfactants used to disperse SWCNTs were explored. Deoxyribonucleic acid (DNA) has been utilized to wrap around SWCNT and disperse them individually in solution (Zheng et al. 2003), while sodium cholate (SC) is a biocompatible surfactant which disperses SWCNT well in solution (Wenseleers et al. 2004; Arnold et al. 2006).

Herein, DNA-and SC-dispersed SWCNT are characterized by near-infrared (NIR) spectroscopy and their toxicity to *C. reinhardtii* is determined.

5.2 Methods

5.2.1 SWCNT Preparation

Single-wall carbon nanotubes dispersed by either DNA or SC (DNA:SWCNT or SC:SWCNT) preparations were made similar to as previously described (Zheng et al. 2003; Lahiji et al. 2007). Briefly, 2 mg CoMoCAT SWCNT of mixed chirality (Sigma-Aldrich; St. Louis, MO) were mixed with 2 mg salmon testes genomic DNA or SC in 2 mL 1X phosphate-buffered saline (PBS). This mixture was sonicated on ice for two hours with a Virsonic XL2020 ultrasonic liquid processor (Misonix; Farmingdale, NY) equipped with a 3.2 mm microtip at approximately 20% power. The resulting solution was centrifuged at 16,000Xg for 90 minutes to remove insoluble SWCNT. For DNA:SWCNT samples, the solution was then filtered through a 100,000 molecular weight centrifugal filter (Millipore; Billerica, MA) three times at 4000 RPM for 15 minutes each to remove DNA that did not associate with SWCNTs. The recovered DNA:SWCNT solution was brought to a volume of 500 μ L with 1X PBS, while SC:SWCNT samples remained in 2 mL. The concentration of the DNA: or SC:SWCNT samples were obtained by gravimetric analysis of 10 μ L of the solution weighed on an Ohaus Discovery Microbalance (Ohaus; Parsippany, NJ).

5.2.2 SWCNT Characterization

Absorption and NIR fluorescence measurements were performed essentially as previously described (Bachilo et al. 2002; Weisman 2010). Briefly, each SWCNT preparation was diluted to a 1% solution with Milli-Q water (Millipore; Billerica, MA). This solution was added to a 10 mm path length cuvette (Sterna Cells; Atascadero, CA) and analyzed on a

NanoSpectralyzer 1 (NS1) (Applied NanoFluorescence; Houston, TX). The absorbance spectrum was collected from 400 - 1600 nm. Additionally, integrated fluorescent emission spectra were obtained using excitation wavelengths of 638 nm, 690 nm, and 784 nm. The resulting spectra were analyzed with ANFSOft (Applied NanoFluorescence; Houston, TX) and prepared in Origin 8 (OriginLab Corporation; Northampton, MA). Fluorescence efficiency measurements, which are fluorescence power collected across the emission spectrum for a single excitation wavelength divided by the absorbance spectrum, are reported as a relative measure of SWCNT dispersion in solution.

5.2.3 Algae Culture

Chlamydomonas reinhardtii wild-type strain cc-1690 was obtained from the Chlamydomonas Resource Center (St Paul, MN). Received algae were initially plated onto 1.5% Sueoka's high salt medium (HSM) agar prepared as previously described (Sueoka 1960). This was used to seed small liquid cultures of HSM for outgrowth, which was kept in a continuous culture of 100 mL HSM. Every 1-2 weeks, the existing culture was diluted to 10% in HSM to allow continuous outgrowth. Culture conditions were sustained at 22-25°C with continuous ambient light and daily mixing.

5.2.4 SWCNT Treatments

Treatments of *C. reinhardtii* in liquid culture with SWCNT preparations or controls were performed in a total volume of 10 mL in 50 mL conical tubes with loosened lids. At day zero, 9 mL HSM was seeded with 1 mL stock algal culture and the appropriate treatment was added. Treatments were: 0, 0.1, 1, 2, 10, or 100 µg/mL DNA:SWCNT or SC:SWCNT, and 100 µg/mL sonicated DNA or SC. Contamination and absorbance controls consisted of 10 mL of just HSM

or HSM with 1 $\mu\text{g/mL}$ DNA:SWCNT or SC:SWCNT. Each treatment and control was performed in triplicate and grown for ten days at the above-noted culture conditions.

5.2.5 Growth Measurements

Every 24 hours, from day one to day 10, each sample was mixed and 300 μL was removed. The optical density of this solution was measured at 750 nm with a Synergy 2 microplate reader and the software Gen5 1.06 (Biotek US; Winooski, VT). The optical density at 750 nm was also measured for prepared absorbance controls of HSM with each individual treatment. The optical density of algae was determined by subtracting control from sample optical density. On the tenth day, optical density of Chlorophyll β and α at 663 nm and 645 nm respectively was obtained. A portion of each sample was centrifuged and resuspended in methanol. This was centrifuged, and optical density of chlorophyll extracted in the liquid phase was measured at 663 nm and 645 nm as previously described (Chochois et al. 2010). At the end of treatments, algae were placed onto microscope slides with a cover slip and imaged using a Leica ATC 2000 light microscope (Leica Microsystems; Wetzlar, Germany) and a TCA 5.0 microscope camera (Ample Scientific; Norcross, GA).

5.3 Results and Discussion

5.3.1 SWCNT Preparation and Characterization

Single-wall carbon nanotubes were dispersed in solution with either DNA or SC. As determined by gravimetric analysis, concentrations of prepared SWCNT solutions ranged from 10 $\mu\text{g}/\mu\text{L}$ to 30 $\mu\text{g}/\mu\text{L}$. Absorbance and NIR fluorescence spectroscopy was then performed on dilutions of each sample to determine SWCNT chirality composition, diameter, and dispersion in solution (Figures 5.1a and 5.1b). The DNA:SWCNT showed an average SWCNT diameter of 0.83 nm, ranging from 0.6 to 1.4 nm (Figure 5.1c). Based on an integrated fluorescence

emission spectrum, the sample of mixed chirality was composed of largely (6,5), (8,4), and (7,5) armchair SWCNT (Figure 5.1e). The SC:SWCNT sample had an average of 1.248 nm, with SWCNT being mostly between 1.2 and 1.3 nm (Figure 5.1d). The integrated fluorescence emission spectrum of this sample showed a majority of SWCNT being of (11,7) armchair chirality (Figure 5.1f). Additionally, efficiency of fluorescence output of the DNA:SWCNT was much higher than SC:SWCNT, which were negative values (Figures 5.1g and 5.1h). NIR fluorescence is only emitted from monodisperse SWCNT suggesting DNA is more effective at dispersing individual SWCNTs than SC (Zheng et al. 2003; Wenseleers et al. 2004; Liu et al. 2009). DNA gives more monodisperse SWCNT by wrapping around the SWCNT in a helical manner masking the hydrophobic surface of the SWCNT. In contrast, SC forms micelles which disperse SWCNT as bundles. Therefore, fluorescence efficiency and SWCNT diameter of each SWCNT sample confirms that DNA:SWCNT are more well-dispersed than SC:SWCNT.

5.3.2 Effects of SWCNT and Dispersion Method on Algal Growth

Chlamydomonas reinhardtii were grown in liquid culture with either DNA:SWCNT, SC:SWCNT, or controls and the culture optical density was measured for ten consecutive days. Results show no normalized optical density for either the high salt medium or 1 µg/mL DNA:SWCNT only controls, meaning no contamination of media or SWCNT sample was present (Figure 5.2). From culture concentrations of 0.1 µg/mL to 100 µg/mL, there are no effects of DNA:SWCNT on optical density of the algae. For all cultures, a plateau in algal growth was reached in the sixth day. There is also no difference with treatment of 100 µg/mL sonicated DNA only.

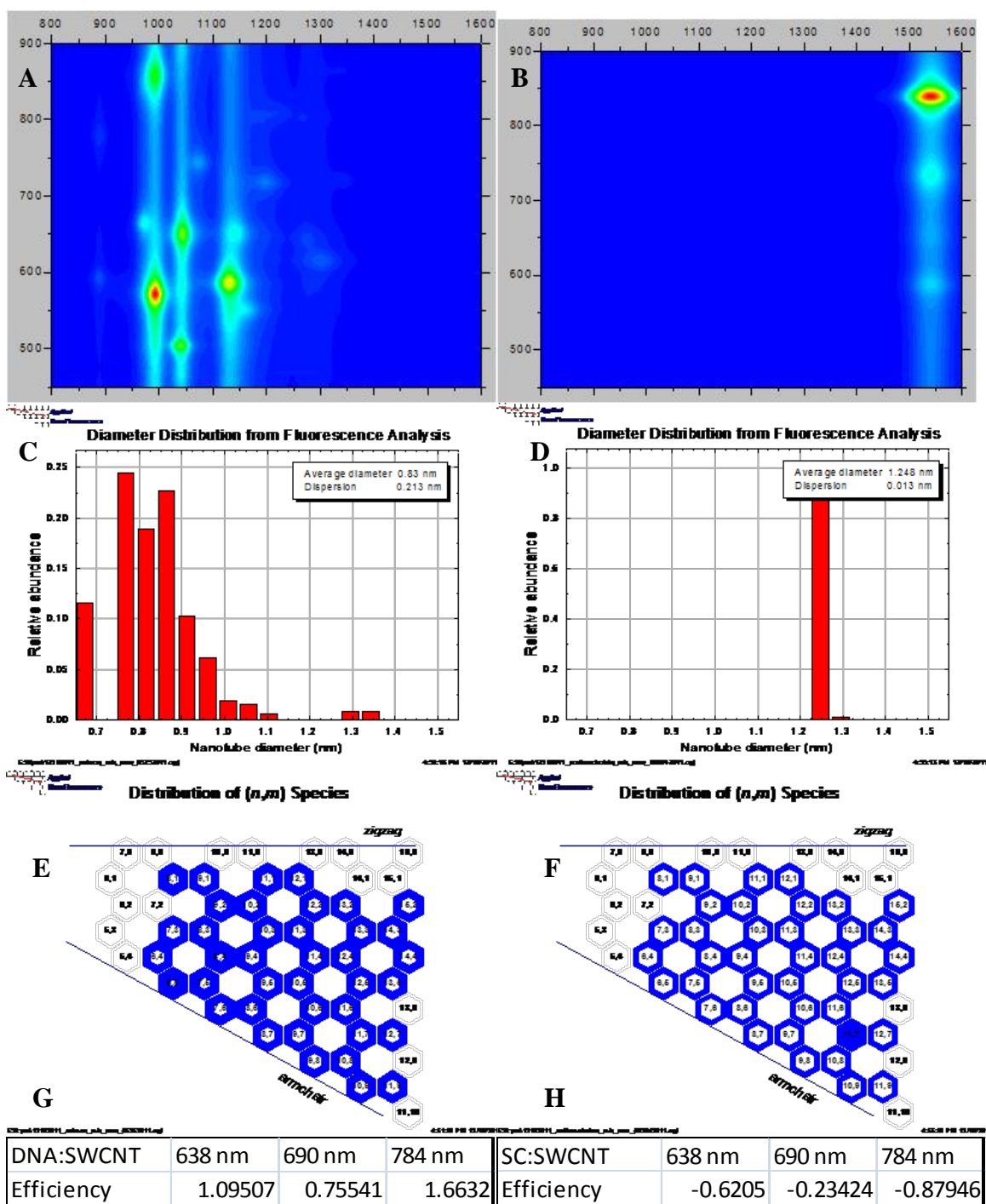


Figure 5.1. Characterization of DNA:SWCNT and SC:SWCNT. (a) and (b) Synthesized fluorescence intensity plot of DNA:SWCNT and SC:SWCNT output respectively. (c) and (d) Calculated density distribution of DNA:SWCNT and SC:SWCNT respectively. (e) and (f) SWCNT chirality distribution of DNA:SWCNT and SC:SWCNT samples respectively. (g) and (h) Fluorescence efficiency measurements at excitation wavelengths of 638, 690, and 784 nm of DNA:SWCNT and SC:SWCNT respectively. Higher numbers represent better dispersion.

Results of algae grown in the presence of SC:SWCNT also show no difference in algal growth optical density at 750 nm due to SC:SWCNT treatment (Figure 5.3). A plateau in growth is evident beginning at day four. There is an initial difference in growth of algae treated with just 100 $\mu\text{g/mL}$ SC at the plateau, which subsides by the ninth day. At the ninth and tenth day, there is an increase in growth of algae treated with 0.1 $\mu\text{g/mL}$, however this is not significant. Growth of SC only-treated algae plateaus at day 6; however standard deviations also overlap with the control at these days suggesting no true difference.

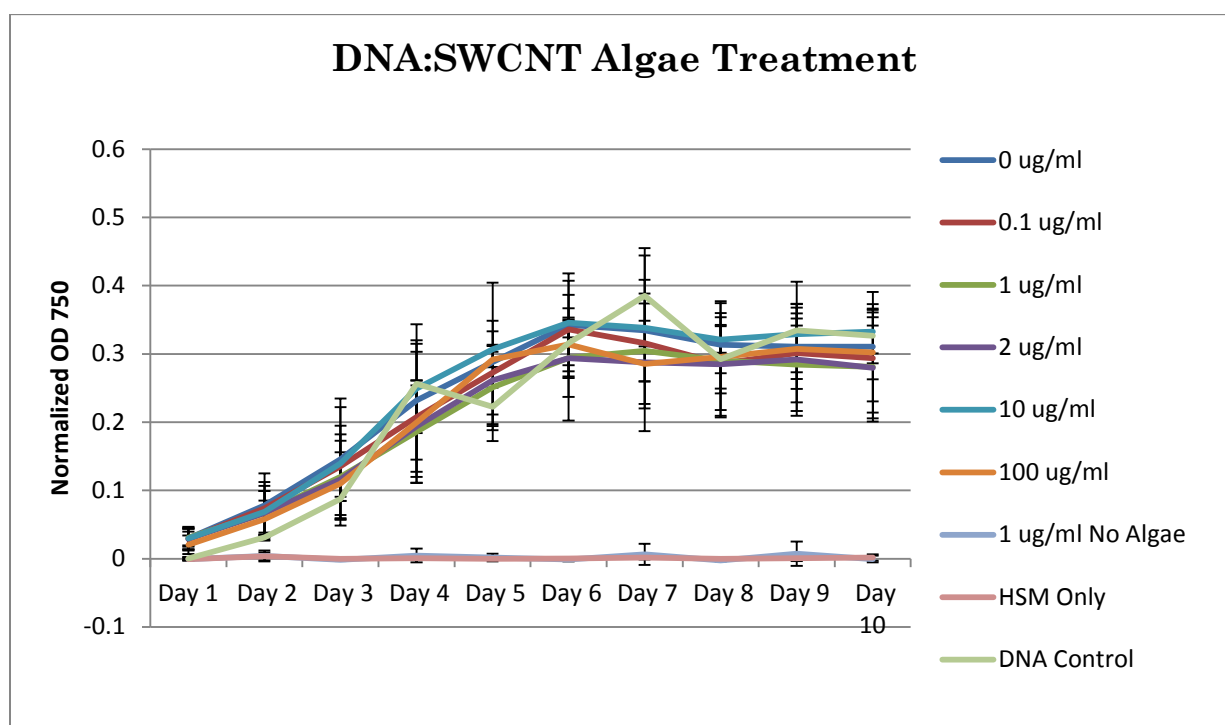


Figure 5.2. Ten day growth curve of *Chlamydomonas reinhardtii* with DNA:SWCNT treatments or controls. Treatments were 0, 0.1, 1, 2, 10, or 100 $\mu\text{g/mL}$, DNA:SWCNT only control, media only control (HSM), or 100 $\mu\text{g/mL}$ DNA control. Errors bars for each point represent (+/-) standard deviation.

After the tenth day, extraction of chlorophyll and absorbance measurements at 663 nm and 645 nm gave chlorophyll content measurements of the algal cultures. At 663 nm, absorbance of chlorophyll β showed no difference between either DNA:SWCNT or SC:SWCNT and their control (Figure 5.4). Additionally, there is no difference between DNA and SC

dispersed SWCNT at any treatment concentration. There is a decrease in SC-only treated algae, however. This may be due to some mitigation by SWCNT being complexed with SC.

Chlorophyll α absorbance at 645 nm showed similar results (Figure 5.5). There is no difference between controls and treatments of DNA:SWCNT and SC:SWCNT at any treatment level.

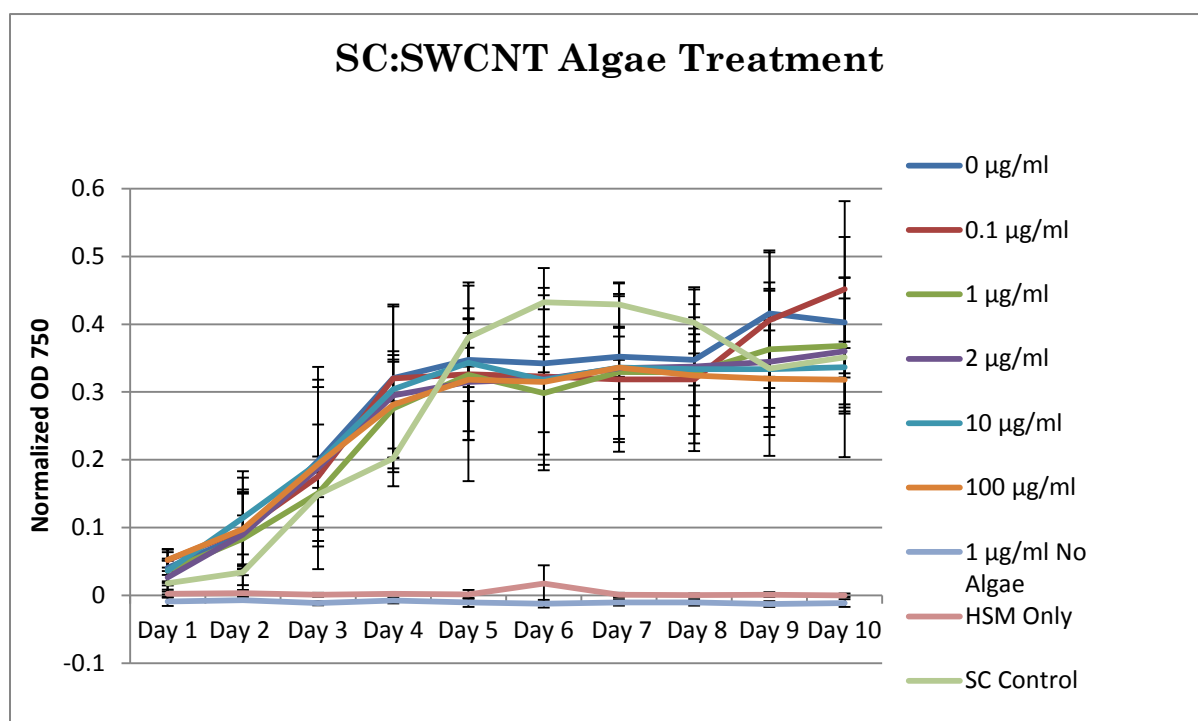


Figure 5.3. Ten day growth curve of *Chlamydomonas reinhardtii* with SC:SWCNT treatments or controls. Treatments were 0, 0.1, 1, 2, 10, or 100 $\mu\text{g/mL}$, SC:SWCNT only control, media only control (HSM), or 100 $\mu\text{g/mL}$ SC control. Errors bars for each point represent (+/-) standard deviation.

There seems to be no difference in *Chlamydomonas reinhardtii* growth or chlorophyll output for DNA:SWCNT or SC:SWCNT treatment relative to control. This is in opposition to previously-reported data showing negative growth effects of SWCNT. Previous findings using 2 $\mu\text{g/mL}$ SWCNT with no surfactant showed a negative effect on growth of algae up to five days (Matorin et al. 2010). This report also showed clear evidence of algal association with SWCNT agglomerations. Non-dispersed SWCNT have also shown to be toxic to bacteria (Kang et al. 2009). Furthermore, a LC_{50} of 2.4 $\mu\text{g/mL}$ has been obtained with *Daphnia magna*. This level

and almost 50 times greater caused no growth effects in the current study. While the stagnant growth conditions used in this study were not conducive to using non-dispersed SWCNT due to hydrophobic precipitation, the previously-reported studies are a comparative benchmark for their toxic effects. The use of well-dispersed SWCNT is environmentally relevant as agglomerations are known to associate with organic matter and become more well-dispersed over time (Lou et al. 2011; Schierz et al. 2012). The extensive studies reported herein suggest there is no negative effect of well-dispersed SWCNT on algal growth.

The reported toxic effect of SWCNT may be due to agglomeration and shading, which was mitigated in this study with more effective dispersion of the SWCNTs. It has been reported that SWCNT toxic effects on the green algae *Pseudokirchneriella subcapitata* were lost with higher concentrations of the surfactant Gum Arabic (Youn et al. 2011). Another study specifically investigating the effects of agglomeration and shading found that SWCNT partially dispersed by natural organic matter had EC₅₀ values of 1.8 µg/mL and 20 µg/mL on the algae *Chlorella vulgaris* and *P. subcapitata* (Schwab et al. 2011). That work suggests that toxic effects may be due to indirect effects on local growing conditions of algae within SWCNT agglomerations, such as competition for resources and reduced light availability.

In the current study, agglomeration and shading by SWCNT did not seem to be as prevalent. No agglomeration of DNA:SWCNT samples were seen either in culture or in micrographs (Figure 5.6). At the highest level of SC:SWCNT treatment, 100 µg/mL, there was some visible agglomeration in culture and under the microscope, but this was not extensive nor was it found at any lower levels. However, microscopic examination shows only a small portion of algae are agglomerated with SC:SWCNT, not nearly as prevalent as with the *P. subcapitata* study. Furthermore, growth at the 100 µg/mL treatment level was not affected.

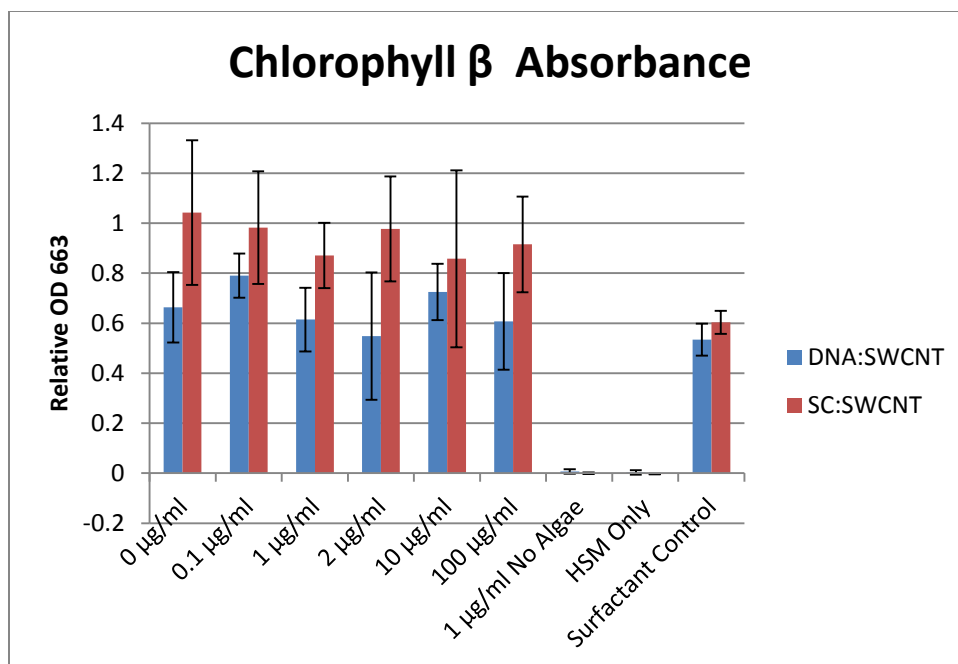


Figure 5.4. Extracted Chlorophyll β content at OD 663 for DNA:SWCNT and SC:SWCNT treatments. Treatments are 0, 0.1, 1, 2, 10, or 100 $\mu\text{g/mL}$, DNA:SWCNT or SC:SWCNT only

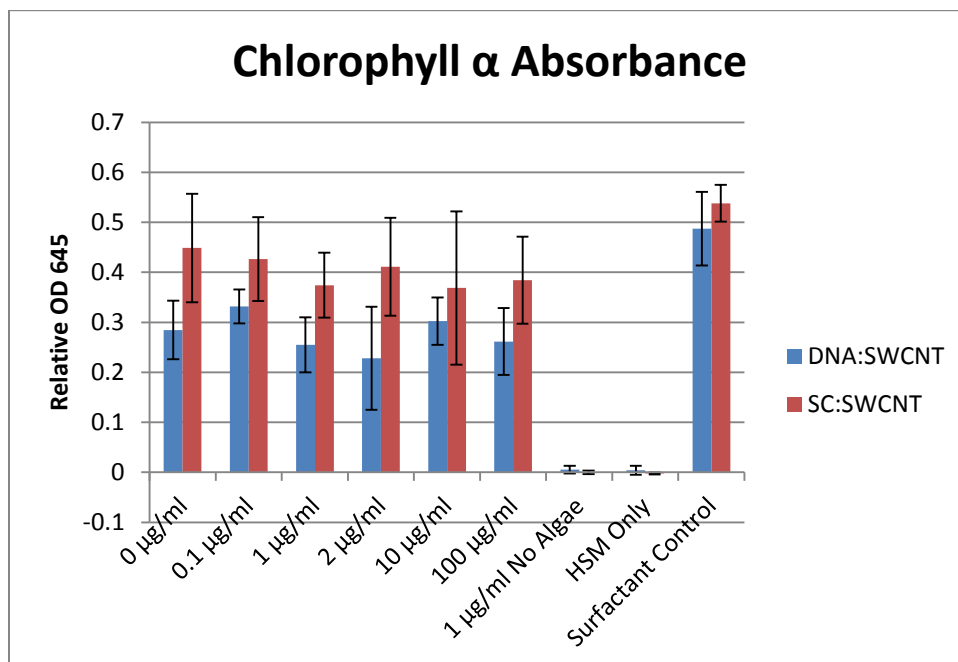


Figure 5.5. Extracted Chlorophyll α content at OD 645 for DNA:SWCNT and SC:SWCNT treatments. Treatments are 0, 0.1, 1, 2, 10, or 100 $\mu\text{g/mL}$, DNA:SWCNT or SC:SWCNT only control, media only control (HSM), or 100 $\mu\text{g/mL}$ DNA or SC control. Error bars for each point represent (+/-) standard deviation.

Additionally, the chosen strain can grow in the presence of nitrogen in the dark and therefore may not be affected by SWCNT shading effects. Results of chlorophyll content measurements however show no difference due to SC:SWCNT at any level. Future work may determine possible effects on chlorophyll efficiency due to agglomeration and shading. Hence, the lack of toxic SWCNT effects is likely due to the well-dispersed nature of DNA:SWCNT and SC:SWCNT. Indeed, work with SWCNT well-dispersed in various surfactants on rat liver cells and human lung fibroblasts showed no toxic effects (Alpatova et al. 2010; Wang et al. 2010). Furthermore, DNA:SWCNT and SC:SWCNT showed no toxic effect on human astrocytoma cells (Dong et al. 2008). A lack in difference of growth between DNA:SWCNT and SC:SWCNT, even though DNA:SWCNT are more well-dispersed, may be because SC:SWCNT do not agglomerate to the point of causing a wide-scale effect on growth.

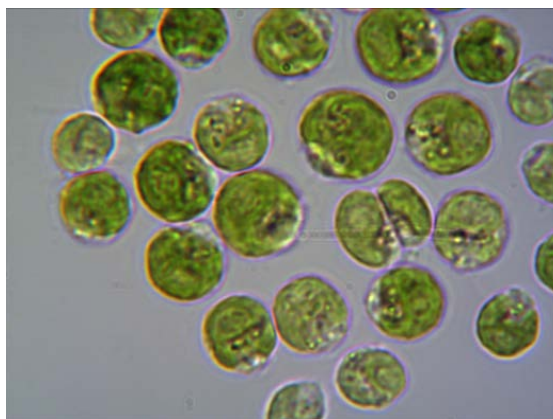


Figure 5.6. Micrograph of algae grown with SWCNT. Representative image of *Chlamydomonas reinhardtii* after ten days in culture with 100 $\mu\text{g/mL}$ DNA:SWCNT. Algae appear healthy and no SWCNT agglomerations are visible.

A detractor from this hypothesis is a study showing that SWCNT act as 'nano darts'. In this, well-dispersed SC:SWCNT effectively punctured bacterial membranes (Liu et al. 2009). This work states that softer bacterial membranes were more susceptible to puncturing by SWCNT, and therefore may not be a factor with the prominent glycoprotein cell wall of *C. reinhardtii* (Harris 2001). It was also noted that a higher rate of bacterial death was obtained by

faster shaking speeds with SWCNT. This was not a factor here as cultures were only mixed once a day. Future studies may be directed to investigate increased shaking speeds of DNA:SWCNT and SC:SWCNT treated cultures. The presented work does, however, agree with other toxicological studies using DNA:SWCNT and SC:SWCNT to treat human cells (Dong et al. 2008). It is therefore unlikely that SWCNT punctured the cell wall of the algae or had any other negative effects. Future work may further investigate the nature of the interaction seen in SC:SWCNT agglomerations and investigate the ability of monodisperse SWCNT to penetrate algal cell walls.

5.4 Conclusions

This study presents findings that SWCNT well-dispersed with DNA or SC does not have an effect on growth or chlorophyll content of *Chlamydomonas reinhardtii* at concentrations of 0.1-100 µg/mL. This is in opposition to previous work that has shown SWCNT can be toxic to algae, bacteria, or other model organisms. However, this supports other research that suggests there is no toxic effect of well-dispersed SWCNT on model organisms or mammalian cells. Thus, increasingly widespread use of SWCNT may have toxic environmental impacts if the materials are poorly dispersed in solution. However, well-dispersed SWCNT, such as those with DNA or SC, may have minimal environmental consequences.

Appendix A

***In Vitro* Selection of Antibody Fragment Molecular Recognition Elements for Prostate Cancer Cells**

A.1 Introduction

Prostate cancer is the most-diagnosed non-skin cancer in the United States, with an estimated 241,740 new diagnoses in 2012 alone (Siegel et al. 2012). It is the second leading cause of cancer-related deaths among males in the U.S., with an estimated 28,170 mortalities in 2012 (Siegel et al. 2012). Approximately one in six U.S. men will get prostate cancer in their lifetime, with one in 30 dying from the disease (Ries et al. 1997). It is therefore important to develop new diagnostic and therapeutic approaches for clinical prostate cancer treatment.

Currently, prostate cancer diagnosis has shifted away from the standard of care: prostate specific antigen (PSA) blood levels. Initially discovered in 1979, serum PSA levels were shown to be higher in men with prostate cancer than those with normal prostates (Wang et al. 1979; Kuriyama et al. 1980). After U.S. Food & Drug Administration approval for prostate cancer detection in 1986, there was an initial increase in number of prostate cancer-related deaths, largely due to more diagnoses (Catalona et al. 1991; Baade et al. 2004). This mortality rate began to decrease due to better awareness, however, as PSA had become the standard bearer in cancer biomarkers (Baade et al. 2004).

In spite of this promise, PSA testing is no longer recommended for use by the United States Preventative Services Task Force (USPSTF) (Chou et al. 2011; Lin et al. 2011; Moyer 2012). This is because serum PSA levels are raised not only due to cancerous prostates, but due to benign prostatic conditions such as prostatic intraepithelial neoplasia (PIN), benign prostate hyperplasia (BPH), and prostatitis (Woolf 1995). Indeed, studies have shown the specificity of the PSA test in prostate cancer diagnosis to be just 24%, meaning there is a 76% over-diagnosis rate (Thompson et al. 2006). Calculations have shown that PSA screening will prevent just one prostate cancer-related death in greater than 1000 men (Moyer 2012). For that small number is

does help, it is estimated to cause many unnecessary biopsies, treatments, and individual stress (Carlsson et al. 2012; Moyer 2012). For that reason, the USPSTF officially gave it a grade ‘D’ in 2012; however some primary care physicians continue to use the test due to its high sensitivity (Barry 2009; Moyer 2012; Pollack et al. 2012).

The promise of PSA, despite its pitfalls, has led to a great deal of research in identifying other diagnostic mechanisms. There have been many biomarkers identified that may have some diagnostic potential (Herness and Naz 2001; Williams and Naz 2010). Many of these, however, do not differentiate between prostate cancer and other prostatic diseases, similar to PSA. Others are expressed in other tissues and are not useful for identification of prostate cancer. The most-studied among these is prostate-specific membrane antigen (PSMA), which has entered into the clinic as a diagnostic and therapeutic tool (Horoszewicz et al. 1987). PSMA, however, is expressed in other non-prostatic normal and malignant tissues (Maraj et al. 1998; Renneberg et al. 1999). Thus, clinical prostate cancer diagnosis relies on digital rectal exams, which can miss tumors, relies on diagnosis after palpable tumors have formed, and misses more prostate tumors than PSA screening (Catalona et al. 1994). Additionally, biopsies are performed, which are invasive and have inherent infection risks (Rodriguez and Terris 1998). Therefore, it is necessary to develop a minimally invasive diagnostic molecular test for the early and specific detection of prostate cancer.

In addition to non-specific diagnostics, current treatments for prostate cancer are partially effective, but have many unwanted side effects. Early-stage, localized prostate cancer is typically either managed with watchful waiting or treated with prostatectomy, radiation therapy, androgen deprivation, cryotherapy, or ultrasonography (Chou et al. 2011). Long-term studies have found that radical prostatectomy confers a significant decrease in mortality compared to

watchful waiting, from 21% to 15% in one trial (Wong et al. 2006; Albertsen et al. 2007; Bill-Axelson et al. 2008; Bill-Axelson et al. 2011). In total, radiotherapy, including external beam radiation therapy and internal brachytherapy, significantly reduces prostate cancer-related mortality compared to no treatment (Tewari et al. 2007; Zhou et al. 2009; Ladgevardi et al. 2010; Stattin et al. 2010). Conversely, androgen deprivation therapy is actually associated with increased risk of mortality, however one study found a slight decrease after a seven year followup period (Lu-Yao et al. 2008; Zhou et al. 2009).

Each of the localized prostate cancer treatment modalities is associated with multiple side-effects. These include urinary incontinence, erectile dysfunction, and quality of life, among others (Chou et al. 2011). In fact, depending on the treatment, a 14-33% higher proportion of men, compared to watchful waiting, suffered from erectile dysfunction and a 20% higher proportion of men had urinary incontinence (Chou et al. 2011). For prostatectomy, operation-related mortality and cardiac events are possible (Yao and Lu-Yao 1999). Additionally, recurrence is possible with all of these treatments. This can be in a more aggressive, androgen-independent form, or other hard-to-treat forms (Feldman and Feldman 2001). For many non-localized cancers, chemotherapy is also used. This is in spite of the fact that many advanced, hormone-refractory prostate cancers are difficult to treat with chemotherapeutics (Yagoda and Petrylak 2006). For prostate cancer, historically 5-fluorouracil (5-FU) was used and had fewer side effects compared to those used today (Yagoda and Petrylak 2006). Currently, docetaxel is the first-line chemotherapeutic, with mitoxantrone sometimes being used in that role based on side effects (Pienta 2001; Pienta and Smith 2005). These are often used with combinations of other agents as well (Pienta and Smith 2005). However, whether alone or in combination, these cause many side effects due to non-specific action. A 70% response rate has been shown for

patients on docetaxel, however as many as 70% of all patients also experienced neutropenia (Extra et al. 1993). Other main side effects include thrombosis, fatigue, hyperglycemia, nausea, vomiting, and others (Pienta and Smith 2005). Clearly, these drugs can be effective, however more specific therapeutics would be more effective and have fewer side effects.

It is possible to utilize molecular targeting to diagnose and specifically deliver therapeutics to prostate cancer. Identification of a Molecular Recognition Element (MRE) is achieved through the process of Selective Evolution of Ligands by EXponential Enrichment (SELEX), also known as *in vitro* selection (Ellington and Szostak 1990; Tuerk and Gold 1990). This process utilizes a large library of random molecules which are screened for their ability to bind to a target of interest. Those that do not bind to the target are removed, and those that do are amplified. Amplified molecules are then subjected to closely-related negative targets to which binding is not desired. Molecules that do not bind to this target are amplified, completing a round of selection. Rounds continue until one or a few strongly-binding and specific molecules are selected. This process has been adapted for use in identifying molecules that bind to whole-cell targets (Morris et al. 1998; Guo et al. 2008). It has also been adapted to incorporate a protein library displayed on various microbiological host surfaces, including yeast (Huse et al. 1989; Gunneriusson et al. 1996; Boder and Wittrup 1997). It is therefore possible to combine these methodologies to identify protein MREs from a yeast-displayed library which bind to the surface of prostate cancer cells.

The work described here utilized a novel variation of SELEX called Decoy-SELEX. This scheme enriches the initial library for molecules which bind to the target of interest similar to SELEX. It focuses, however, on stringent negative selections using targets to which the MRE should not bind. This work has identified MREs which bind to a prostate cancer cell line, and

not to various benign prostate cells lines. It utilized a human non-immune single-chain Fragment variable (scFv) library displayed on the surface of *Saccharomyces cerevisiae*. The obtained MREs will be useful for both specific diagnosis and treatment of prostate cancer.

A.2 Methods

The Decoy-SELEX process was used in order to obtain a prostate cancer cell-specific MRE (Figure A.1). A library of non-immune human single-chain Fragment variable (scFv) antibody fragments was displayed on the surface of yeast. Seven rounds of selection were completed, enriching for those scFvs which bound to the androgen-dependent prostate cancer cell line LNCaP and subtracting those that bound to the benign prostate cell lines HGPIN, BPH-1, and BHPRE1, as well as the protein PSMA (Table A.1).

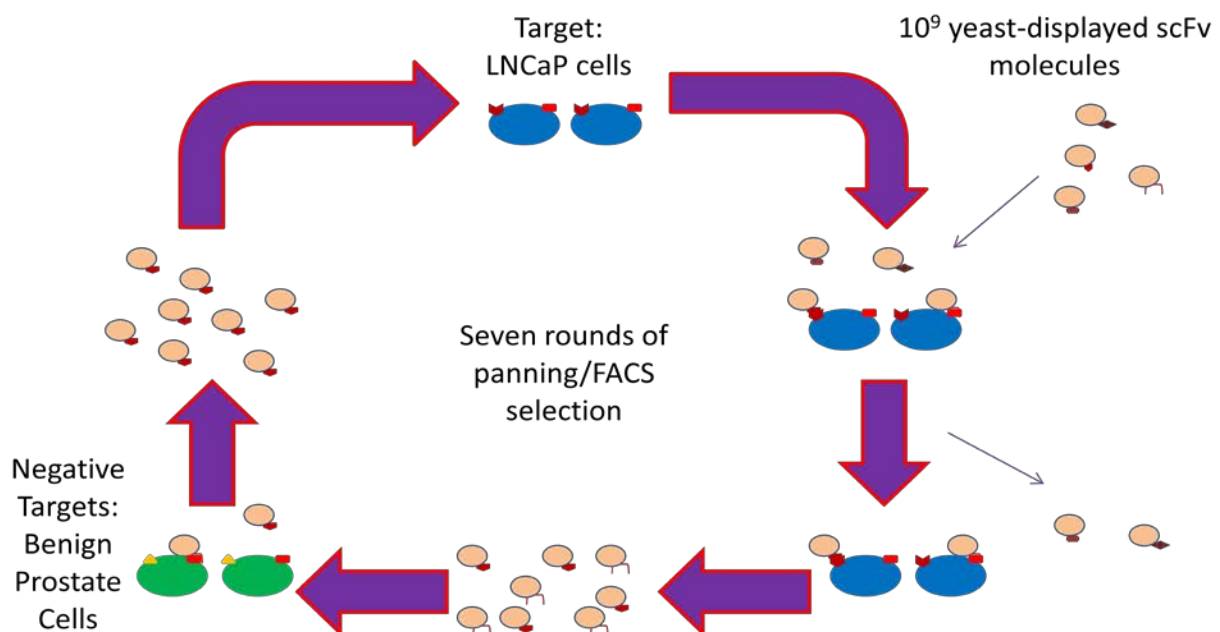


Figure A.1. Scheme depicting the Decoy-SELEX process. A library of 10^9 yeast-displayed scFv molecules are incubated with the target LNCaP cell line. Those that bind are amplified and incubated with negative targets. Those that do not bind the negative targets are amplified and subjected to additional rounds of *in vitro* selection.

A.2.1 Cell Culture and Materials

In order to obtain a prostate cancer cell-specific MRE, prostatic cell lines were used. For general maintenance, each line was passaged every 5-7 days in a T75 cell culture dish with media changed every 2-3 days. The cells were grown in a 37°C incubator with 5% carbon dioxide and humidity. The androgen-dependent prostate cancer cell line LNCaP was obtained from the American Type Culture Collection (ATCC) (Manassas, VA) and cultured in RPMI 1640 with L-Glutamine and 25 mM HEPES (Cellgro; Manassas, VA) and 10% Fetal Bovine Serum (FBS) (Fisher Scientific; Pittsburgh, PA) and 1X antibiotic/antimycotic mixture (ab/am) (Cellgro) (Horoszewicz et al. 1983). The HPV-18-immortalized High Grade Prostatic Intraepithelial Neoplasia (HGPIN) cell line was a generous gift from Dr. Mark Stearns (Drexel University; Philadelphia, PA) and was cultured in Defined KSFM (Gibco; Grand Island, NY) with 5% FBS and 1X ab/am (Wang et al. 1999). The SV40T-immortalized Benign Prostate Hyperplasia (BPH-1) cell line was a generous gift from Dr. Simon Hayward (Vanderbilt University; Nashville, TN) and was cultured in RPMI-1640 with L-Glutamine and 25 mM HEPES and 10% FBS and 1X ab/am (Hayward et al. 1995). The spontaneously-immortalized intermediate prostate stem cell line BHPRe1 was also obtained as a generous gift from Dr. Simon Hayward (Vanderbilt University) and cultured in DMEM/F12 (Cellgro) supplemented with 5% FBS, 1X ab/am, 1% insulin/transferrin/selenium (Gibco), 0.4% bovine pituitary extract (Sigma; St. Louis, MO), 5 ng/mL epidermal growth factor (Gemini Bio-Products; West Sacramento, CA), and 1X ab/am (Jiang et al. 2009). The androgen-independent DU-145 prostate cancer cell line was obtained from ATCC and cultured in EMEM (Cellgro) with 10% FBS and 1X ab/am (Stone et al. 1978). The androgen-independent prostate cancer cell line PC-3 was also obtained from ATCC and cultured in F12K media (Cellgro) with 10% FBS and

Table A.1. Decoy-SELEX scheme for identification of prostate cancer cell-specific MREs.

Selection Round	Target	Yeast Number	% Yeast Expression	Human Cell Number	Incubation Time	Volume	Separation Method
1 (+)	LNCaP	1 X 10 ¹⁰	34.2	~8 X 10 ⁶	3 hours	15 mL	Panning
1 (-)	HGPIN	1 X 10 ¹⁰	38.5	~7 X 10 ⁶	30 minutes	15 mL	Panning
	BPH-1			~9 X 10 ⁶	30 minutes		
	BHPrE1			~7.5 X 10 ⁶	30 minutes		
2 (+)	LNCaP	1 X 10 ¹⁰	7.04	~9 X 10 ⁶	2 hours	15 mL	Panning
2 (-)	HGPIN	1 X 10 ¹⁰	6.34	~8.5 X 10 ⁶	1 hour	15 mL	Panning
	BPH-1			~9.5 X 10 ⁶	1 hour		
	BHPrE1			~8 X 10 ⁶	1 hour		
3 (+)	LNCaP	1 X 10 ¹⁰	8.85	~9 X 10 ⁶	1 hour	15 mL	Panning
3(-)	HGPIN	1 X 10 ¹⁰	6.7	~7 X 10 ⁶	2 hours	15 mL	Panning
	BPH-1			~9 X 10 ⁶	2 hours		
	BHPrE1			~8 X 10 ⁶	2 hours		
4 (+)	LNCaP	1 X 10 ⁷	29.65	1 X 10 ⁶	30 minutes	2.5 mL	FACS
4 (-) a	HGPIN	1 X 10 ⁷	6.31	1 X 10 ⁶	30 minutes	1.5 mL	FACS
4 (-) b	BPH-1	1 X 10 ⁷	7.65	1 X 10 ⁶	30 minutes	2 mL	FACS
4 (-) c	BHPrE1	1 X 10 ⁷	4.78	1 X 10 ⁶	30 minutes	1.5 mL	FACS
4 (-) d	PSMA*	1 X 10 ⁷	17.3	1 nmol protein	30 minutes	1 mL	FACS
5 (+)	LNCaP	1 X 10 ⁷	24.02 (Lo)/21.72 (Hi)	1 X 10 ⁶	30 minutes	5 mL	FACS
6 (+)	LNCaP	1 X 10 ⁷	21.2/19.36	1 X 10 ⁵	30 minutes	2.75 mL	FACS
6 (-) a	HGPIN	1 X 10 ⁷	16.73/13.5	1 X 10 ⁶	30 minutes	2.5 mL	FACS
6(-) b	BPH-1	1 X 10 ⁷	7.86/11.92	1 X 10 ⁶	30 minutes	2.5 mL	FACS
6(-) c	BHPrE1	1 X 10 ⁷	41.42/48.85	1 X 10 ⁶	30 minutes	2.5 mL	FACS
6(-) d	PSMA*	1 X 10 ⁷	37.26/8.53	1 nmol protein	30 minutes	2 mL	FACS
7 (+)	LNCaP	1 X 10 ⁷	25.67/15.63	1 X 10 ³	30 minutes	2.5 mL	FACS

*PSMA is prostate-specific membrane antigen.

1X ab/am (Kaighn et al. 1979). The HPV-18-immortalized normal prostatic epithelium cell line RWPE-1 was obtained from ATCC and cultured in Defined KSFM (Gibco) plus 1X ab/am

(Bello et al. 1997). The spontaneously-immortalized early prostate stem cell line NHPrE1 was a generous gift from Dr. Simon Hayward (Vanderbilt University) and cultured in the same media as BHPPrE1 (Jiang et al. 2009).

A.2.2 scFv Library and Growth

A human non-immune scFv library displayed on the surface of *Saccharomyces cerevisiae* with 10^9 diversity was a gift from Dr. Dane Wittrup (Massachusetts Institute of Technology; Cambridge, MA) (Boder and Wittrup 1997; Feldhaus et al. 2003). The library had a full diversity of at least 10^9 antibody fragments and has the ability to be serially propagated without loss of diversity (Feldhaus et al. 2003). The full library was cloned into the pPNL6 plasmid in the EBY100 yeast strain. It was originally amplified in standard dextrose media with casamino acids (SD+CAA), which contains 0.5% casamino acids, 2% dextrose, 0.17% yeast nitrogen base without ammonium sulfate amino acids, 0.53% ammonium sulfate, 1.019% sodium dihydrogen phosphate, 0.856% sodium monophosphate, and supplemented with 10 U/mL penicillin, 10 μ g/mL streptomycin, and 80 μ g/mL ampicillin that had been vacuum filtered through a 0.22 μ m filter. The library was amplified to an OD of approximately 1.5 at 30°C with shaking at 250 RPM. Surface expression of the library was achieved by expressing the scFv under control of a galactose promoter in standard galactose plus casamino acids media (SG+CAA), which substituted 2% galactose, 2% raffinose, and 0.1% dextrose for the dextrose in SD+CAA. An initial input of 0.5 OD was amplified to approximately 1.5 OD with shaking at 250 RPM at 20°C. After each induction growth period, surface expression of the library was confirmed by flow cytometry analysis. The monoclonal anti-HA tag antibody clone 16B12 conjugated to either DyLight 488 (Columbia Biosciences; Columbia, MD) or AlexaFluor 488 (Invitrogen; Grand Island, NY) was used to stain scFv-expressing yeast in yeast wash buffer (YWB)

consisting of phosphate-buffered saline (PBS), 0.5% bovine serum albumin, and 2 mM EDTA. The samples were run on either a Cell Lab Quanta SC (Beckman Coulter; Brea, CA) or a FACSCalibur (BD Biosciences; San Jose, CA) flow cytometer equipped with a 488 nm argon laser and 525 nm emission filter. At least 5% partial induction of the surface library was confirmed and deemed sufficient for selection incubations.

A.2.3 In Vitro Selection

Seven rounds of Decoy-SELEX were performed in order to obtain a scFv MRE specific for the LNCaP prostate cancer cell line. The first three rounds of selection were performed by panning and the last four by fluorescence-activated cell sorting (FACS). The first three rounds of panning-based selection were performed directly in a T75 cell culture flask. For Round 1(+) selection, LNCaP cells were grown to 80-90% confluence and the media was removed. The cells were gently washed with calcium- and magnesium-free phosphate-buffered saline (PBS), which was removed. The cells were then incubated with 10^{10} yeast to ensure total representation of the library diversity. The naïve yeast library was amplified as discussed previously and scFv surface expression was induced. After surface expression confirmation, the library was suspended in 15 mL YWB. The library was placed into the flask containing LNCaP cells and placed on a 37°C shaker at 25 RPM for three hours. After incubation, yeast not bound to LNCaP cells were removed, and the LNCaP cells were gently washed three times with 15 mL YWB. It was then noticed, however, that LNCaP cells were less than 25% confluent in the flask, likely due to EDTA metal chelation and subsequent cell removal.

To determine optimal buffer conditions, separate 90% confluent LNCaP T75 flasks were incubated with the following six buffers at 37°C with shaking at 25 RPM for three hours: 1) YWB, 2) YWB with no EDTA, 3) YWB with no EDTA and 0.1% FBS, 4) YWB with no EDTA

and 0.5% FBS, 5) YWB with no EDTA and 1% FBS, and 6) YWB with no EDTA and 10% FBS. After incubation, remaining attached cells were visually estimated and compared to before incubation. Buffer 1 caused less than 25% of LNCaP to be attached, while Buffers 2-4 resulted in approximately 50% attachment, and Buffers 5 and 6 caused almost no detachment (90% confluent LNCaP cells), likely due to the FBS supplement. Therefore, Buffer 5 was chosen as an appropriate incubation buffer and named yeast selection buffer (YSB).

After determination of an appropriate incubation buffer, the Round 1(+) incubation was performed as described above, however with expressing yeast suspended in YSB. After three washes, 100 mL SD+CAA was added to the flask to allow for amplification of yeast bound to LNCaP cells. This was grown overnight, and this enriched library was prepared for a round of negative selection. For Round 1(-) selection, scFv-expressing yeast were suspended in YSB and incubated with rinsed HGPIN cells at 80-90% confluence for 30 minutes at 37°C with shaking at 25 RPM. The supernatant containing yeast-displayed scFvs not bound to HGPIN cells was removed and added to rinsed BPH-1 cells under the same conditions, removed again and added to rinsed BHPRE1 cells under the same conditions. The serially-incubated supernatant was centrifuged to obtain yeast, which were suspended in SD+CAA for amplification. Two more rounds of panning-based selection were performed in this manner, with decreasing incubation times for positive rounds and increasing incubation times for negative rounds.

The ensuing four rounds of selection were performed using FACS-based sorting. For Round 4(+) selection, LNCaP cells grown to 80-90% confluence were fluorescently dyed with CFSE (Invitrogen) according to manufacturer's instructions. They were then dissociated from the flask with CellStripper reagent (Cellgro) to prevent cell surface protein digestion associated with trypsinization. Cells were then suspended in YSB and counted with a Scepter equipped

with 60 μm sensors (Millipore; Billerica, MA). In Round 4, all yeast were fluorescently dyed with Syto61 (Invitrogen) according to manufacturer's instructions. A total of 10^7 yeast were suspended in YSB and mixed with 10^6 LNCaP in a volume of approximately 2.5 mL. They were mixed by inversion for 30 minutes at 37°C and placed on ice before FACS sorting. The sample was then sorted with a FACS Aria (BD Biosciences), with excitation at 488 nm from a sapphire solid state laser and 633 nm from a HeNe laser and 525 nm and 650 nm emission filters. Events identified as bound yeast and LNCaP cells were collected. In Round 4(+), there were two separate populations of bound cells which were collected and amplified separately. These were called the 'Hi' and 'Lo' populations and kept separate through the following rounds of selection and subjected to the same growth and selection conditions.

In Round 4(-)a, for each of the Hi and Lo populations, yeast were prepared for selection in the same manner and incubated with 10^6 HGPIN cells dyed the same with inversion for 30 minutes at 37°C in YSB. The sample was subjected to FACS and events that indicated yeast that were not bound to HGPIN cells were collected and amplified. Those were then prepared for selection and incubated with 10^6 dyed BPH-1 cells for Round 4(-)b and subjected to FACS with unbound yeast collected and amplified. The enriched library was then prepared for Round 4(-)c and subjected to FACS after incubation with 10^6 dyed BHPRE1 cells. Unbound yeast were collected and amplified, then prepared for selection and incubated with 1 nmol recombinant full-length PSMA protein, which is expressed in non-prostatic normal and tumor tissue, as well as normal prostate cells (Silver et al. 1997) (Abnova; Walnut, CA), pre-incubated with AlexaFluor 488-conjugated anti-GST tag antibody (Invitrogen). The yeast which showed single fluorescence corresponding to Syto61 were kept and amplified.

After post-Round 4 amplification, yeast from the Hi and Lo populations were prepared for Round 5(+) selection. They underwent the same process as before, however they were fluorescently labeled with anti-HA monoclonal antibody 16B12 conjugated to AlexaFluor 647 (Invitrogen). This was done for the following rounds to ensure only strongly-expressing yeast were collected. For Round 5(+) selection, 10^7 yeast were incubated with 10^6 LNCaP cells. The yeast bound to LNCaP cells were collected, amplified, and prepared for Round 6(+) selection. A total of 10^5 dyed LNCaP cells were used to increase stringency. Round 6(-) was performed similar to Round 4(-) with the exception of antibody-labeled yeast to increase stringency. Unbound yeast were collected, amplified, and prepared for Round 7(+). In that round, 10^3 yeast were used to increase stringency and select the scFvs with highest affinity.

A.2.4 Fluorescent Imaging of scFv Library Enrichment

To visualize enrichment of the scFv library after 7 rounds of selection, fluorescent micrographs were obtained. LNCaP cells were grown in a 6-well culture dish with number 1.5 microscopic cover glasses to ~90% confluency and fluorescently labeled with CFSE as described above. Yeast from the Round 0, post-Round 7(+) Lo, and post-Round 7(+) Hi libraries were cultured as described above. Surface expression of the scFv was confirmed by antibody labeling and flow cytometry as described. 10^7 yeast were then labeled with anti-HA antibody conjugated to AlexaFluor 647 (Invitrogen) and incubated with LNCaP cells in 1 mL YSB in the 6-well dish. Cells were incubated at 37°C for 30 minutes with shaking at 25 RPM. The supernatant was removed and the cells were washed three times with 2 mL YSB. The cells were then fixed with 10% formalin (Azer Scientific; Morgantown, PA) for 5 minutes and washed three times with 1 mL PBS then three times with 1 mL double-distilled water. Cover slips were dried and flipped onto a microscope slide prepared with 20 uL Fluorogel (Electron Microscopy Sciences; Hatfield,

PA), and dried. Mounted cells were imaged on a Zeiss LSM 510 Confocal Microscope (Zeiss; Thornwood, NY) using an Argon laser at 488 nm and a HeNe laser at 688 nm for excitation with emission filters at 505-530 nm band pass and 650 nm. Images were processed with Zeiss LSM Image Browser (Zeiss).

A.2.5 Sequencing of scFv Library

A representative sample of scFv-genes from the naïve library (Round 0) was sequenced to determine diversity of the library. Additionally, this was completed for the enriched post-Round 2(-) and post-Round 3(-) libraries. It was also performed for each of the Hi and Lo populations following Round 4(-), Round 6(-) and Round 7(+) selections. To do this, yeast amplified from the previous round of selection were plated onto 2% agar plates containing 1% yeast extract, 2% peptone, and 2% dextrose (YPD). Yeast were grown for 36-48 hours at 30°C, and individual colonies were chosen for polymerase chain reaction (PCR) amplification.

Yeast colonies were picked and placed into double distilled water and boiled, which served as template for the PCR reaction. The reaction ingredients were as follows: 400 nM forward (5'-GTACGAGCTAAAAGTACAGTG-3') and reverse (5'-TAGATACCCATACGACGTTC-3') pPNL6 primers (Eurofins MWG Operon), 250 µM deoxynucleotide triphosphates, 5% dimethyl sulfoxide, 1X Phusion Reaction Buffer (New England Biolabs, Ipswich, MA), 2 units Phusion High-Fidelity DNA polymerase (New England Biolabs), and double distilled water to 100 µL. Reactions conditions were: initial denaturation at 98°C for 30 seconds; 35 cycles of 98°C for 10 seconds, 50°C for 30 seconds, and 72°C for 30 seconds; and final extension at 72°C for 10 minutes. Results were analyzed using agarose gel electrophoresis and those PCRs that contained bands corresponding to the scFv gene were prepared for sequencing. They were purified using a PCR purification kit (IBI Scientific; Peosta,

IA) and sent for DNA sequencing (Eurofins MWG Operon; Huntsville, AL) using both the forward and reverse pPNL6 primers. In total, at least 30 sequences were obtained for each enriched library noted above. Analysis was then performed by translating the DNA sequence to protein using the ExPASy translate tool (Swiss Institute of Bioinformatics; <http://web.expasy.org/translate/>). Hemagglutinin, c-Myc, and linker protein tag sequences, landmarks of the scFv expression scaffold, were identified to ensure sequence quality (Boder and Wittrup 1997). Sequences were compared for similarities and duplicates within an enriched library were identified to determine diversity of the library.

A.2.6 Secretion and Purification of Selected scFvs

From the Rounds 6(-) and 7(+) sequences, three scFvs were chosen for further study due to their abundance in the enriched libraries: HiR6.8, HiR7.6, and HiR7.8. Sequences were PCR amplified with the same ingredients and under the same conditions as above; however pPNL9 primers were used. For full length scFv sequences (HiR6.8), the forward pPNL9 (5'-GACGTTCCAGACTACGCTGGTGGTGGTGGTTCTGCTA-3') and reverse pPNL9 (5'-GGGTTAGGGATAGGCTTACCCTGTTGTTCTAGAATTCCG-3') were used. For truncated scFv sequences (HiR7.6 and HiR7.8) containing only the heavy chain, forward pPNL9 and modified reverse pPNL9 (5'-GGGTTAGGGATAGGCTTACCGAACTCTGAAGAGACGGTGACC-3') were used. The PCR product was analyzed by agarose gel electrophoresis and the desired scFv-encoding gene was excised from the gel. DNA was purified from the gel using the GFX gel purification kit (GE Healthcare; Piscataway, NJ). Purified scFv-encoding DNA was then used for transformation of scFv-secreting yeast.

To express and secrete the scFv, the encoding gene was cloned into the pPNL9 plasmid which was transformed into the YVH10 strain yeast, both generous gifts from Dr. Dane Wittrup (Massachusetts Institute of Technology). Prior to cloning, the pPNL9 plasmid was double digested with the restriction enzymes NotI and SfiI (New England Biolabs) and analyzed by agarose gel electrophoresis. The plasmid was prepared for further studies by phenol/chloroform extraction and ethanol precipitation. Transformation was performed by gap repair cloning and the LiAc TRAFO method as previously described (Gietz and Woods 2002). Briefly, YVH10 yeast with no plasmid were grown in YPD broth. A fresh sample was taken and centrifuged, then resuspended in the following: 67% of 50% polyethylene glycol 3350, 0.1 M lithium acetate, 13.9% of 2 mg/mL boiled carrier DNA (salmon testes genomic DNA; Sigma), approximately 1 µg purified pPNL9-amplified scFv DNA, and approximately 0.3 µg double-digested pPNL9 plasmid, with water to 360 µL. The yeast were incubated at 42°C for 1.5 hours, centrifuged, and suspended in water. They were then plated on selective SD+CAA agar with 0.8% tryptophan (Trp) and grown at 30°C for 48-96 hours until individual colonies formed.

Individual colonies were chosen and subjected to colony PCR amplification as previously described, with the appropriate pPNL9 primer set used. The purified PCR product was sent for DNA sequencing to confirm the insert. The colony was then amplified in selective SD+CAA+Trp media overnight at 30°C with shaking at 250 RPM. Approximately 5×10^8 yeast were then centrifuged and suspended in 50 mL expression media containing 1% yeast extract, 2% peptone, 2% galactose, 2% raffinose, and 0.1% dextrose, plus antibiotics as listed above (YPG/R). Yeast were grown for approximately 72 hours with shaking at 250 RPM at 20°C for secretion of scFv.

After scFv secretion, supernatant was removed from the yeast by centrifugation. The supernatant was concentrated with a 9 kDa centrifugal filter (Pierce). The concentrated scFv solution was then brought to a volume of 10 mL with PBS, 0.5 M NaCl, and 0.05% Tween 20. Then, 200 μ L Ni-NTA agarose resin (Qiagen; Valencia, CA) was added and mixed with inversion at 4°C for 2 hours for purification of 6-His tagged scFvs. The mixture was centrifuged and supernatant removed and washed three times with 10 mL PBS, 0.5 M NaCl, and 0.05% Tween 20 solution. The scFv was then eluted from the agarose resin three times with 250 μ L of 200 mM imidazole in PBS. Pre-incubation, wash, and elution fractions were then concentrated using a 3 kDa molecular weight centrifugal filter (Pierce).

Concentrated fractions were then analyzed using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE). Samples were mixed with loading dye, denatured and loaded into a 4-20% Tris-HEPES gel. Samples were run and the gel was stained with Coomassie blue staining buffer. The gel was imaged, in which proper secretion and purification was confirmed by obtaining bands of the appropriate size for each scFv. Remaining elution fractions were combined, with extra imidazole removed with 3 kDa centrifugal columns. The concentration of the scFv was determined using spectroscopy with a NanoDrop (Thermo Scientific; West Palm Beach, FL). The scFv was then diluted to a working concentration of 10 μ M in PBS.

A.2.7 Affinity Binding Assays with scFvs

In order to determine the binding affinity of each secreted scFv with the LNCaP cell line, saturation binding assays were performed with flow cytometry with few modifications to previous work (Benedict et al. 1997; Boder et al. 2000). LNCaP cells were grown to 80-90% confluency and removed from the flask with CellStripper reagent. The cells were counted with a Scepter equipped with 60 μ m sensors. Then, 2×10^5 cells were placed into a 500 μ L total

volume of YSB. LNCaP cells incubated with no antibody served as a negative fluorescent control, and LNCaP cells incubated with 1 μ L anti-PSMA antibody conjugated to AlexaFluor 488 (BioLegend; San Diego, CA) served as a positive fluorescent control. For saturation binding assays, the appropriate amount of each scFv was incubated on ice for 30 minutes with 2 μ L of the monoclonal anti-HA antibody clone 16B12 conjugated to DyLight 488 (Columbia Biosciences). Concentrations of 0, 1, 10, 100, 1000, 10000, 50000, 100000, 150000, 200000 pM of the scFv were used. The pre-incubated scFv and secondary antibody were mixed with LNCaP cells and placed on a rotisserie at 37°C for 30 minutes. After this period, the cells were centrifuged for 7 minutes at 300xg at 4°C. The supernatant was removed and the cells were resuspended in 500 μ L YSB.

Samples were run on a FACSCalibur (BD Biosciences) as previously described. The mean fluorescence intensity (MFI) for each sample was recorded. The MFI for the 0 pM incubation, which had only the secondary fluorescent antibody, served as background fluorescence and was subtracted from the MFI of each incubation. These data were graphed using Origin 8 software (OriginLab Corporation; Northampton, MA). The data were fit with a nonlinear regression model of single-event binding given by the equation, $Y = ((B_{\max} * X) / (K_d + X)) + NS * X$, where B_{\max} is maximum binding, K_d is the dissociation constant, and NS is nonspecific binding. Binding assays for HiR6.8 and HiR7.8 were performed in triplicate, with average dissociation constants calculated and standard errors derived from the equation. Binding assays for HiR7.6 will be performed in the future in triplicate.

A.2.8 Specificity Binding Assays of scFvs

In order to determine the specificity of MRE binding, cross-binding assays were performed with the cell lines used as negative targets in the selection. LNCaP, HGPIN, BPH-1,

and BHPRE1 cells were grown to 80-90% confluency and removed from the flask with CellStripper reagent. Cells were then counted with a Scepter using 60 μm sensors and 2×10^5 cells were placed into a final volume of 500 μL YSB. For each cell line, cells with no antibody served as a negative control. Concentrations of 0 or 100000 pM scFv were incubated with the secondary anti-HA AlexaFluor 488 antibody for 30 minutes on ice. Each of these were then mixed with each cell line and incubated on a rotisserie at 37°C for 30 minutes. The cells were then centrifuged, the supernatant removed, and they were resuspended in 500 μL YSB. Each sample was then run on a FACSCalibur as previously described and the MFI recorded. The MFI of the 0 pM incubation, which had only the secondary fluorescent antibody, served as background fluorescent labeling and was subtracted from the 100000 pM incubation. Data were normalized to LNCaP binding, which was set to 100%, and graphed to show comparative binding between the cell lines. HiR6.8 and HiR7.8 assays have been performed in triplicate, with average values calculated and Student's T-tests performed to determine statistical significance. HiR7.6 will be assayed in the future in triplicate.

In order to determine binding to other prostatic cell lines used not used in the selection, cross-binding assays were performed. LNCaP, the androgen insensitive prostate cancer cell lines DU-145 and PC-3, and the normal prostatic cell lines RWPE-1 and NHPRE1 were used for this experiment. The cells were prepared as noted for the other cross-binding assays and incubated with the scFv as described above. Samples were run on a FACSCalibur and the MFI was normalized and graphed as noted. HiR6.8 and HiR7.8 have been performed in triplicated, with average values calculated and Student's T-tests performed to determine statistical significance. HiR7.6 will be assayed in the future in triplicate.

A.2.9 Fluorescent Imaging of scFv Binding

In order to visualize binding of each scFv, fluorescent images were taken. LNCaP cells were grown in a 6-well culture dish with number 1.5 microscopic cover glasses to ~90% confluency. For each scFv, a 100 nM concentration was pre-incubated with anti-HA antibody conjugated to DyLight 488 as described for binding assays, in addition to LNCaP cells with anti-PSMA antibody, and those with secondary antibody only in a total volume of 500 μ L YSB. Cells were incubated at 37°C for 30 minutes with shaking at 25 RPM. The supernatant was removed and the cells were washed once with 2 mL YSB. The cells were then fixed with 10% formalin (Azer Scientific; Morgantown, PA) for 5 minutes and washed three times with 1 mL PBS then three times with 1 mL double-distilled water. Cover slips were dried and flipped onto a microscope slide prepared with 20 μ L Fluorogel (Electron Microscopy Sciences; Hatfield, PA), and dried. Mounted cells were imaged on a Zeiss AxioImager Z2 Fluorescent Microscope (Zeiss; Thornwood, NY) using an Argon laser at 488 nm for excitation with a band pass emission filter at 505-530 nm.

A.3 Results and Discussion

A.3.1 Selection of MREs

In order to obtain prostate cancer cell-specific MREs, seven rounds of the Decoy-SELEX process were completed. Sequencing of the scFv-encoding plasmid in the initial naïve library, as well as after Round 2 and Round 3 of panning selection, was performed. Out of at least 30 sequences obtained, there were no duplicate sequences in any library. Additionally, there were no duplicates between the libraries. This suggests that all three libraries were highly diverse, with no particular sequence having been selected for.

After three rounds, FACS-based selections were performed. In Round 4 (+), there were two separate populations of yeast binding to LNCaP cells. One was characterized by having relatively high Syto61 (yeast) fluorescence with low CFSE (LNCaP) fluorescence, and this was designated the ‘Hi’ population. The other was characterized by having relatively low Syto61 (yeast) fluorescence with high CFSE (LNCaP) fluorescence, and was designed the ‘Lo’ population. These were sorted separately and kept separate through the remaining rounds of selection, while each was subjected to identical growth and selection conditions.

After the negative selections in Round 4, sequencing of the scFv-encoding genes in the library revealed duplicate sequences within individual libraries. The Lo population was comprised of 29% HiR7.8 and between 3% and 13% of 13 other molecules (Figure A.2).

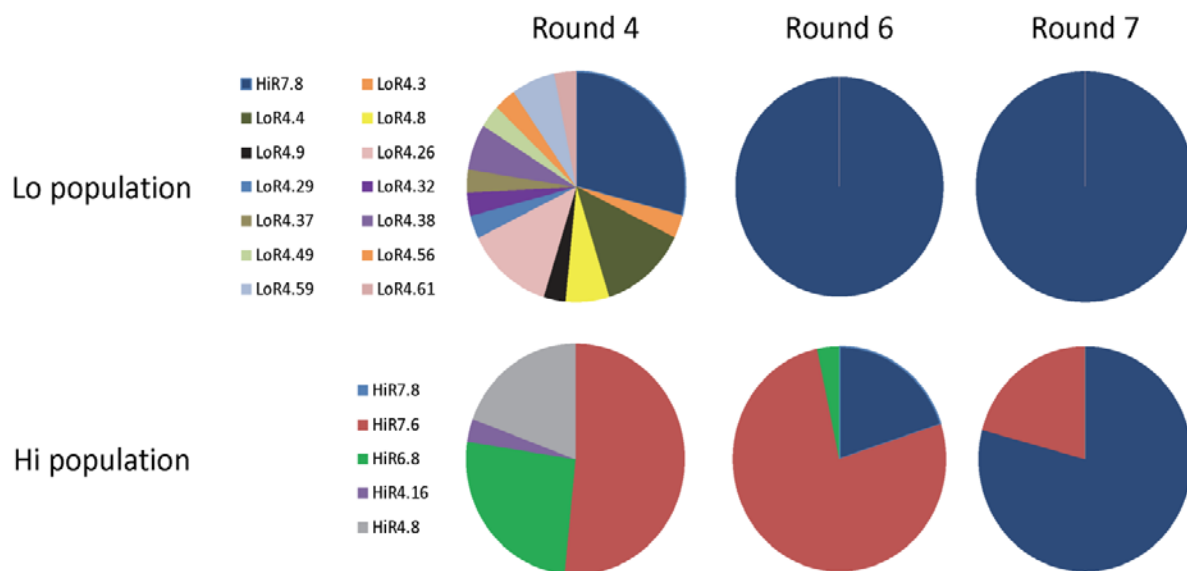


Figure A.2. Enrichment of the Lo and Hi yeast-displayed scFv populations after FACS-based selection rounds. Evolution of library diversity after Round 4, the first round of FACS-based selection, Round 6, and Round 7. Lo population is the top row the Hi population on the bottom. The names of the sequences represented by each color in the pie charts are shown to the left of Round 4.

Interestingly, HiR7.6 and HiR6.8 were not present in the Lo population. In the Hi population, however, HiR7.6 comprised 52% of the population, while HiR6.8 was 26%. One other scFv was 3% and a final molecule was 19%. Interestingly, HiR7.8 did not appear in the Round 4 Hi population. It is also of note that HiR7.6 and HiR7.8 did not appear in any previous library sequencing, however HiR6.8 did appear once in the Round 3 library.

Following the sixth round of selection, the Lo and Hi libraries were sequenced. In the Lo population, 30 out of 30 (100%) of the sequenced molecules were HiR7.8, and there were no longer any other detectable scFvs in the library (Figure A.2). This suggests that, within this population, HiR7.8 was strongly enriched for in Rounds 5 and 6 of selection. In the Hi population, 77% of the library was HiR7.6 while 3% was HiR6.8. Interestingly, HiR7.8 now comprised 20% of the library, and no other molecules were present. This suggests that HiR7.6

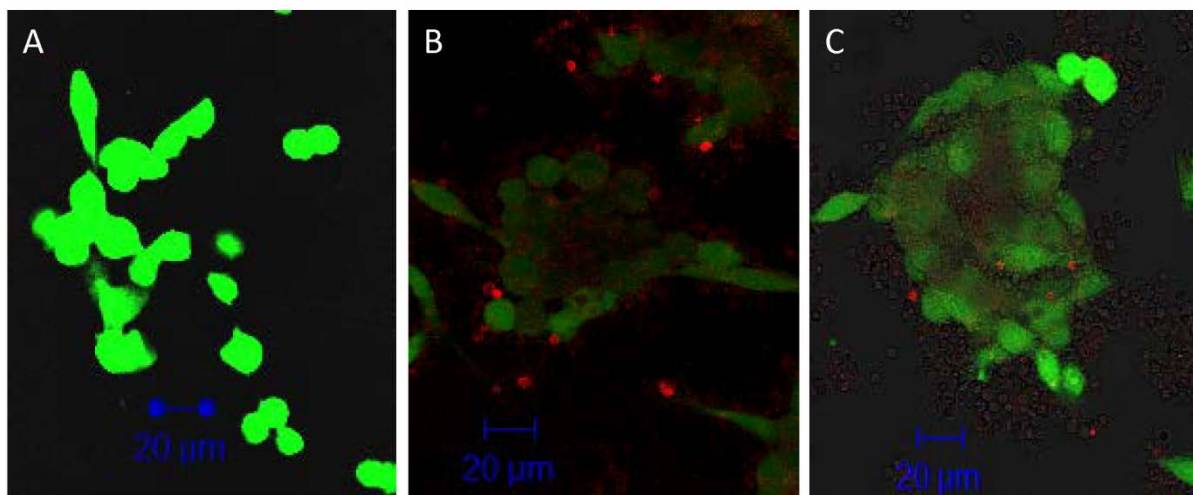


Figure A.3. Fluorescent micrographs of yeast binding enrichment. Confocal microscope images at 200X of LNCaP cells (green) incubated different scFv-displaying yeast libraries. (a) LNCaP incubated with Round 0 library. (b) LNCaP incubated with Round 7(+)Lo library. (c) LNCaP incubated with Round 7(+)Hi library. Images show enrichment of yeast library through seven rounds of selection for yeast-displayed scFv specific for LNCaP cells.

outcompeted HiR6.8 within this population. It also suggests that HiR7.8 was present previously in the library, but not detected, and had outcompeted HiR6.8.

Following a very stringent Round 7(+) selection, using 10,000 fewer target cells than yeast, and 1,000 times fewer LNCaP than Round 6(+), sequencing further revealed the scFvs present in the library. In the Lo population, 100% of the molecules present were again HiR7.8 (Figure A.2). While there may have been other scFvs in the library, it is unlikely after two consecutive rounds of not detecting any besides HiR7.8. In the Hi population, HiR6.8 was no longer detected, while HiR7.6 was now just 16% of the library. HiR7.8 comprised 84% of the library, suggesting it strongly outcompeted HiR7.6. It is possible that if more rounds of selection were completed under more stringent conditions, HiR7.8 would totally comprise the Hi population as well.

Visualization of library enrichment for scFvs binding to LNCaP cells was performed. This provided evidence that few or none of the molecules in the Round 0 library were binding to LNCaP cells (Figure A.3). However, after seven rounds of selection, both the Lo and Hi populations were strongly specific for LNCaP cells (Figure A.3). This, along with the sequence evolution, validates the sequence enrichment capabilities of the novel Decoy-SELEX process. From this obvious enrichment through seven rounds of Decoy-SELEX, HiR7.8 was chosen as the most likely prostate cancer cell-specific MRE and underwent further characterization. However, to determine why it outcompeted HiR6.8 and HiR7.6, those molecules were also chosen for further characterization. It is also important to note that HiR6.8 is a full-length scFv, comprising both a heavy and light chain, with a predicted molecular weight of approximately 30.6 kDa for the scFv itself (Table A.2). HiR7.8 and HiR7.6, however, only contain a heavy chain, as they are truncated just before the chain linker. They have a predicted molecular weight

of approximately 16.0 and 15.6 kDa, respectively, and have 81% sequence identity and 86% sequence similarity. Truncated sequences containing only heavy chains were found in sequencing of the naïve library in the original published description (Feldhaus et al. 2003). This may be relevant *in vivo*, as it has been shown that smaller scFvs have greater tumor penetration ability than larger antibodies (Colcher et al. 1998).

Table A.2. Amino acid sequence of selected prostate cancer cell MREs.

Sequence Name	Sequence*	Predicted Molecular Weight
HiR6.8	DVPDYALQASGGGGSGGGGSGGGGSASQVQLVQSGAEVKKPGASVKVSCKASGY TFTDYMHVWRQAPGQGLEWMGWINPNSGGTNYAQKFQGRVTMTRDTSISTAYM ELSSLRSDDTAIYYCARDADSGSMSAIIYWFNLWGRGTLVTVSSGILGSGGGGS GGGGSGGGGSDIQLTQSPSSLSASVGDRVTITCRASQSI SRFLNWFYQQKPGKAP KLLIYGGSSLSQGVPSRFSGGGSGTDFTLTISLQPEDFATYYCQQSYSKFWTF GQGTKVEIKSGILEQKLISEEDL	30.6 kDa
HiR7.6	DVPDYALQASGGGGSGGGGSGGGGSASQVQLQESGPGLVKPSGTLSTCAVSGG SISSSNWWSVWRQPPGKELEWIGEIHHSGSTNYNPSLKSRTISVDKSKNQFSL KMRSVTAADTAVYYCARVEEWPYDALDMWGQGTMTVTSSEF	15.6 kDa
HiR7.8	DVPDYALQASGGGGSGGGGSGGGGSASQVQLQESGPGLVKPSQTLSTCTVSGD SIYSSGHYWSVWRQHPGKLEWIGYIYASGRTYYNPSLESRTMSVDTSKNQSS LKLTSVTAADTAVYYCARDSDRTWYKAFDTWGQGTMTVTSSEF	16.0 kDa

*Underlined amino acids represent the heavy chain. Double underlined amino acids represent the light chain. HiR7.6 and HiR7.8 do not contain a light chain. Non-underlined amino acids represent scFv structural elements.

A.3.2 Affinity Binding Assays of scFvs

Once HiR6.8, HiR7.6, and HiR7.8 were identified for further characterization, each was cloned into a secretion vector, expressed, and purified. HiR6.8 and HiR7.8 were used in fluorescent saturation binding assays with LNCaP cells as described above, plotted, and fit with nonlinear regression analysis (Figure A.4). HiR6.8 exhibited an equilibrium dissociation constant (K_d) of 33.19 +/- 22.18 nM (Table A.3). The binding affinity of HiR7.6 for LNCaP

cells will be assayed in the future. The binding affinity of HiR7.8 is 27.28 +/- 15.85 nM. The slightly higher affinity of HiR7.8 for LNCaP cells may be a partial explanation as

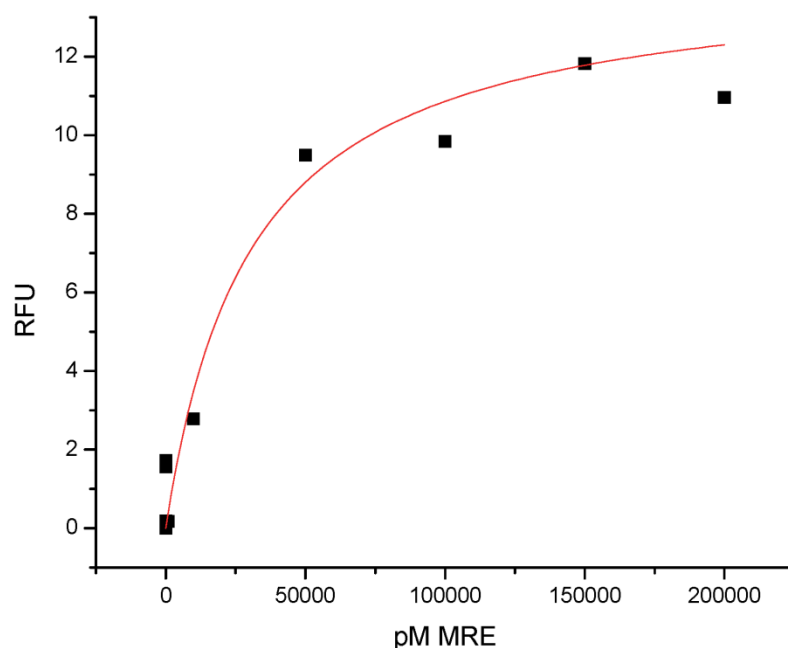


Figure A.4. Plot of fluorescent equilibrium binding assay of HiR6.8 and LNCaP cells. Calculated dissociation constant (K_d) of this assay is 30.52 +/- 27.08 nM.

Table A.3. Equilibrium dissociation constant (K_d) of prostate cancer cell MREs.

HiR6.8	HiR7.6	HiR7.8
33.19 +/- 22.18 nM	*	27.28 +/- 15.85 nM

Data are reported as K_d +/- Standard Error as given by the binding equation. “*” denotes that the data are yet to be acquired.

to why it was enriched for throughout Decoy-SELEX. The dissociation constants for each of these scFvs is in the low-nanomolar range, which is similar to previous reports of scFvs isolated from the selection process (Schier et al. 1996; Benedict et al. 1997; Böldicke et al. 2001). More importantly, it is in a range that has previously been shown to confer strong pharmacokinetic properties (Adams et al. 1998; Jackson et al. 1998). It is possible that further mutagenesis and screening may select for scFvs with higher affinity for the cell surface target (Schier et al. 1996; Boder et al. 2000).

Binding of both HiR6.8 and HiR7.8 to LNCaP cells is further evidenced by fluorescent micrographic analysis. Binding of each scFv is clear when pre-incubated with a secondary fluorescent antibody then with LNCaP cells and imaged for fluorescent staining (Figure A.5). Fluorescence is much brighter than LNCaP cells incubated with fluorescent secondary-only control. Interestingly, both antibodies have a semi-circular or horseshoe-like staining pattern for each cell. Therefore, fluorescence may be visualized in the cytosol surrounding the nucleus. It may be possible that the images represent cellular uptake of the scFv-secondary antibody complex. This has previously shown to be possible and even likely with scFvs alone or in complex with other molecules (Gao et al. 2003; Nielsen et al. 2006; He et al. 2010). This may have an important role in the clinical application of these MREs. The potential ability to use these scFvs as therapeutic targeting agents in the form of antibody-drug conjugates (ADCs) is enhanced by scFv internalization (Schrama et al. 2006). Potential uptake of the scFv by LNCaP cells may be explored in future studies. First, confirming cytoplasmic staining using a nuclear dye such as DAPI would be useful. This could then be furthered by subcellular fractionation experiments. These are potentially interesting further directions stemming from fluorescent imaging of scFv binding.

A.3.3 Specificity Binding Assays of scFvs

Purified HiR6.8 and HiR7.8 were each assayed for their ability to bind to cell lines other than LNCaP. HiR7.6 will be assayed in the future for its binding to other cell lines. These included the negative targets in the selection process, HGPIN, BPH-1, and BHPRe1 (Figure A.6a and Figure A.7a). They also included androgen-independent prostate cancer cell lines DU-145 and PC-3 as well as normal prostate epithelium cell lines RWPE-1 and NHPRe1 (Figure A.6b and Figure A.7b).

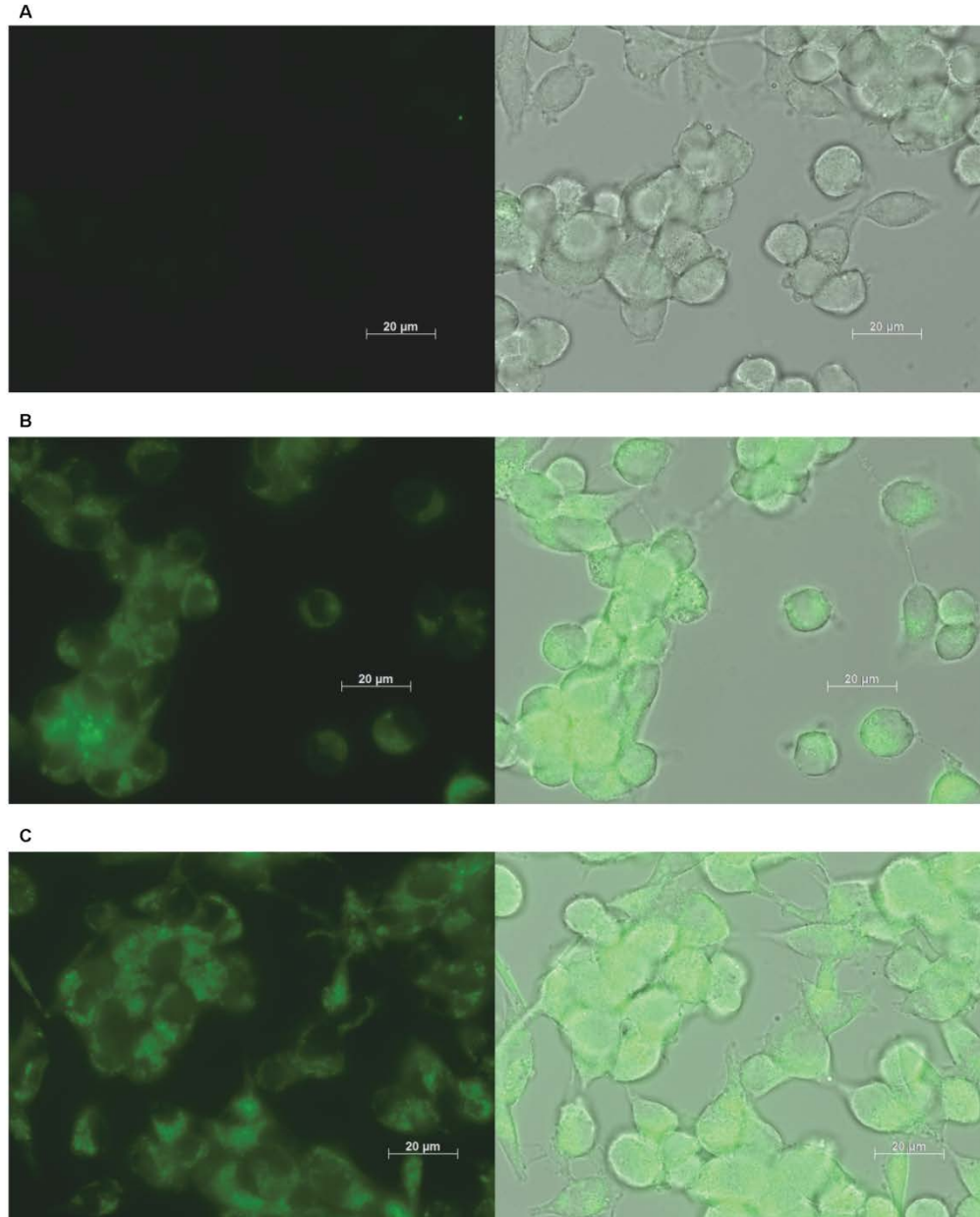


Figure A.5. Micrographs of scFv binding to LNCaP cells. Each panel shows micrographs at 630X of fluorescence from AlexaFluor 488 in green (left) and combined fluorescent plus brightfield image (right). (a) LNCaP cells incubated with only secondary antibody. (b) LNCaP cells incubated with HiR6.8 plus secondary fluorescent antibody. (c) LNCaP cells incubated with HiR7.8 plus secondary fluorescent antibody.

From these experiments, it is clear that HiR6.8 binds to all three negative cell lines as well as the other two androgen-dependent prostate cancer cell lines assayed. Binding to many of these lines was as good as or better than binding to LNCaP cells. This is a likely explanation as

to why HiR6.8 was eventually eliminated from both populations, in spite of its low dissociation constant (Figure A.6). HiR7.8, however, showed excellent selectivity for LNCaP cells (Figure A.7). Binding to HGPIN, BPH-1, and BHPRe1 cells all was much less than to LNCaP, in fact they were negative. This means that the MFI of cells incubated with 100000 pM HiR7.8 and secondary fluorescent antibody was less fluorescent than cells incubated with secondary fluorescent antibody only. This can be explained by pre-incubation and binding of the scFv with the secondary antibody, causing background non-specific binding of the secondary antibody to the cells to be less. Therefore, the negative relative MFI suggests that the antibody is present and binding to the secondary antibody, but not to the cells. Additionally, binding to the four cell lines not used in the selection was either negative or much less relative to binding to LNCaP cells. In sum, this suggests that HiR7.8 is highly specific to LNCaP cells and therefore useful in both the therapeutic targeting and disease detection of prostate cancer.

Biologically, the selectivity of HiR7.8 for LNCaP cells is important. There is a complete absence of binding to all benign cell lines studied, whether they represent normal prostatic cells, BPH, or PIN. This suggests that the cell surface molecule HiR7.8 is binding to is not displayed on the surface of benign prostatic cells. This is supported by previous studies that have identified many genes differentially expressed between the benign conditions represented here and prostate cancer cells (Chakrabarti et al. 2002; Zeng et al. 2004; Chen et al. 2011).

Additionally, there is no binding to the androgen-independent prostate cancer cell lines DU-145 or PC-3. This leads to the conclusion that the cell surface antigen is displayed in the earlier, androgen-dependent stage of prostate cancer. This is possible considering previous studies have found a wide array of genes that are differentially expressed from androgen

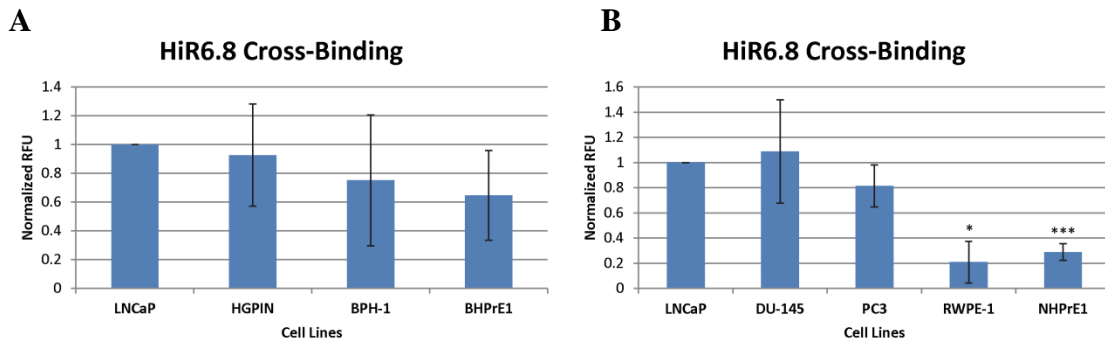


Figure A.6. Cross-binding of HiR6.8 to negative targets and other cell lines. Binding of HiR6.8 to (a) each cell line used in the selection or (b) or prostatic cell lines. Data are normalized to LNCaP binding at 100% for comparison. "*" represents $p < 0.05$, "***" represents $p < 0.001$.

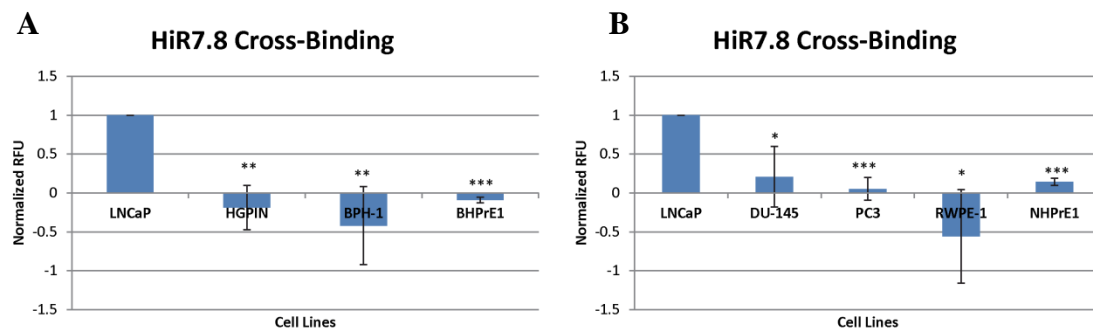


Figure A.7. Cross-Binding of HiR7.8 to negative targets and other cell lines. Binding of HiR7.8 to (a) each cell line used in the selection or (b) other prostatic cell lines. Data are normalized to LNCaP binding at 100% for comparison. "*" represents $p < 0.05$, "***" represents $p < 0.01$, "****" represents $p < 0.001$.

dependent to independent disease progression, many of which are downregulated (Amler et al. 2000; Karan et al. 2002; Chen et al. 2006). In fact, downregulation of expression levels from LNCaP to both DU-145 and PC-3 cells has been found before (Yang et al. 1998; Aalinkeel et al. 2004; Okamura et al. 2008; Liu et al. 2009) It is unclear if HiR7.8 would bind to other androgen-dependent prostate cancer cells, or cells earlier in the progression of the disease, however future work may determine this. Determining binding will be aided by identifying the exact cell surface molecule to which HiR7.8 is binding, as has been done before with other MREs (Robert et al. 2006; Rérole et al. 2011). This type of work has even identified novel cell surface molecules previously unknown to be involved in disease progression (Berezovski et al.

2008). It is therefore possible that this work not only created a novel targeting agent, but also identified a novel target.

HiR6.8, however, is not specific for any one type of cell. This may be because it is binding to some widely-expressed prostate-specific gene such as PSMA (Wright et al. 1995), or because it is binding to any one of a number of common cell surface molecules. Determining the cell surface antigen for HiR6.8 will be important as it may be useful as a broad marker of prostatic cells similar to PSCA or p63 (Signoretti et al. 2000; Tran et al. 2002), or it may be useful in marking a particular cell surface antigen such as previous scFvs for ICAM-1 (Conrad et al. 2009). Therefore, in spite of its spectrum of binding, the high affinity of HiR6.8 for its cell surface target will allow the scFv to be useful for future applications, even if it is not useful in the specific targeting of prostate cancer cells.

The high affinity and specificity of HiR7.8 for LNCaP prostate cancer cells suggest it has potential in future therapeutic and diagnostic applications in prostate cancer. It is unclear whether it will bind to prostate cancer tissue, however it is likely that it will based on binding specificity and affinity similar to previously-selected MREs (Ni et al. 2011). This will rely on surface expression of its cognate antigen. The selective scFv will likely be useful for drug delivery as part of an antibody-drug conjugate (Senter 2009), for chemotherapeutic delivery (Sievers and Linenberger 2001), for radiotherapy delivery (Regino et al. 2009), or possibly even as a therapeutic by itself, depending on the function of the cell surface molecule (Stebbing et al. 2000). Additionally, it will likely have use in specific diagnosis of prostate cancer, whether coupled with an *in vivo* imaging agent (Elsässer-Beile et al. 2009; Veiseh et al. 2010), a capture agent for circulating tumor cells (Nagrath et al. 2007), or as a tissue stain after biopsy (Johnston et al. 1986). Finally, identification of the cell surface markers which these MREs bind to may

possibly identify novel proteins, new functions, or expression patterns that aid in treatment and diagnosis of prostate cancer.

A.4 Conclusions

This work has selected a MRE with high affinity for the androgen-dependent prostate cancer cell line LNCaP. Furthermore, the selected MRE has a high selectivity for the androgen-dependent prostate cancer cell line, and not for benign prostatic cell lines or for androgen-independent prostate cancer cells. These characteristics will allow the selected scFv to be useful for both therapeutic and diagnostic applications in the clinical treatment of prostate cancer. With targeted therapeutics and more specific diagnostics, it is possible that the disease burden of the most-diagnosed cancer among males in the United States will be lessened.

Appendix B

***In Vitro* Selection of Single-Stranded DNA Molecular Recognition Elements for the Pesticide Malathion**

B.1 Introduction

Malathion is the mostly commonly-used organophosphate pesticide in the United States. Approximately 16.7 million pounds are used annually, with 12 million pounds being applied to crops (Wilson et al. 2003). Many other uses include in sewage systems, plant nurseries, home use, and recreational land maintenance (Wilson et al. 2003). Three major historical applications in the U.S. were Boll Weevil eradication, Medfly suppression, and mosquito suppression (Newhart 2006). Its widespread use portends the benefits of malathion usage.

Due to its many uses, malathion has become an environmental contaminant with multiple routes of exposure. Aerial applications often drift to non-target locations, including waterways, up to 200 meters away (LeNoir et al. 1999; Newhart 2006). This method can also lead to volatilization of up to 22.6% of that applied, leading to reduced air quality levels and vaporized contamination spread potentially over large distances (Turner et al. 1991; LeNoir et al. 1999; Ferrari et al. 2003). After one large application, malathion was found in freshwater ponds at levels up to 49.4 µg/L (Segawa et al. 1991). Additionally, runoff due to storms after application has led to levels up to 787.1 µg/L being found (Ando et al. 1996). Detection in groundwater and surface water has also occurred at levels up to 6 µg/L during normal conditions (Newhart 2006). In the state of West Virginia, malathion has been found at levels that exceed guidelines for environmental contamination (Ferrari and Assessment 1997; Ator 1998). Furthermore, one study found that 33.7% of corn grain, 27.8% of cultivated and frozen blueberries, 24.6% of strawberries, and 19.3% of celery samples on the market tested had detectable levels of malathion (USDA 2009). Clearly, release of malathion into the environment and its interaction with humans and other organisms is occurring.

The presence of malathion in the environment is of particular concern to human health. The U.S. Environmental Protection Agency categorizes the pesticide as having "suggestive evidence of carcinogenic potential" (USEPA 2012). This is supported by studies showing induction of mammary epithelial cell proliferation and other carcinogenic effects by malathion through acetylcholinesterase inhibition in rats (Cabello et al. 2001). Additionally, there is a 55%-72% increase in the odds of Attention-Deficit/Hyperactivity Disorder (ADHD) being diagnosed in children that have ten-fold higher malathion exposure levels (Bouchard et al. 2010). Higher concentrations have also been found to have cytotoxic and genotoxic effects on human and other mammalian cells (Giri et al. 2002; Moore et al. 2010). These effects suggest that increased levels of environmental malathion concentrations are detrimental from a public health perspective.

Ecologically, non-target effects of malathion also have severe negative impacts. To various fish species, malathion is highly toxic, with LC_{50} values as low as 0.1 $\mu\text{g/L}$ for brown trout (USGA 2000). Many aquatic invertebrates are also highly susceptible to exposure, with LC_{50} values below environmentally relevant levels, down to 0.5 $\mu\text{g/L}$ (USEPA 2006). It also has other negative effects at sublethal concentrations to a variety of aquatic species (Key et al. 1998; Beauvais et al. 2000; Nguyen and Janssen 2001). Furthermore, the pesticide has been shown to have negative survival effects at environmentally relevant concentrations on tadpoles and frogs (Fordham et al. 2001), small mammals (Newhart 2006), reptiles (Holem et al. 2008), and non-target beneficial insects (Gary and Mussen 1984). It is therefore important that individual species and entire ecosystems are protected from malathion exposure.

Currently, detection of malathion in biological and environmental samples is mostly achieved using gas chromatography (GC). While this method is sensitive and methods for

malathion detection have been improved upon, it is also time- and labor-intensive and requires expensive and expertly-used equipment (Zweig and Devine 1969; Fournier et al. 1978; Wilson et al. 2003). Other methods of malathion exposure include monitoring blood cholinesterase activity or excretion of metabolites (Fenske and Leffingwell 1989; Wilson et al. 2003). There have also been antibody assays developed for malathion detection, however these antibodies have not been assayed for their affinity or only assayed for cross-reactivity to a few of malathion's major chemically-related metabolites (Nishi et al. 2003; Brun et al. 2005; Vera-Avila et al. 2005). As the antibodies were not identified for their selectivity, it is likely that they bind to major metabolites. This is similar to an antibody which binds to a whole class of molecules similar to malathion (Yan et al. 2009). Therefore, it is necessary to identify a molecule that has been identified for its high affinity and specificity for malathion and not its metabolites.

Identification of such a molecule can be accomplished using the Selective Evolution of Ligands by EXponential Enrichment (SELEX). Binding molecules identified with this process are termed Molecular Recognition Elements (MREs). This *in vitro* selection process was originally described by two separate laboratories and MREs are selected from a large library (10^9 - 10^{15}) of different molecules (Ellington and Szostak 1990; Tuerk and Gold 1990). This library is incubated with the target of interest, and those molecules that do not bind to it are removed. Those that do are amplified and can be subjected to negative selections. In those, molecules that do not to the negative targets are amplified. This iterative process continues with increasingly stringent selection conditions until one or a few highly selective molecules with high affinities are identified. A novel variation of this process developed in our laboratory is termed Decoy-SELEX. Using this method, the library is enriched for the target of interest. However, the focus of the selection is on what the MRE should not bind to utilizing stringent

negative selections. Therefore, the Decoy-SELEX method is ideal for identifying MREs with both high affinity and selectivity for the target of interest.

In this work, the Decoy-SELEX method was utilized to identify an MRE for the pesticide malathion. The selection scheme was designed to enriched a single-stranded DNA (ssDNA) library for molecules which bind to malathion. Those molecules which bound to major metabolites of malathion were removed. Those metabolites included dimethyl thiophosphate (DMTP), dimethyl dithiophosphate (DMDTP), malathion dicarboxylic acid (MDA), and malaaxon (Buratti et al. 2005). The MRE was also designed to not bind to other pesticides found in similar amounts and environments as malathion including propanil (Köck et al. 2010), 2,4-D acid (Baker et al. 2000), and atrazine (McCarthy and Fuiman 2008). Additionally, bovine serum albumin (BSA) was used as a negative target as a model for large, globular proteins. The selected MRE may be used as a sensor in a rapid, field-deployable detection device (Soh et al. 2009; Yang et al. 2010; Lin et al. 2011).

B.2 Methods

B.2.1 Decoy-SELEX for Malathion-Specific MREs

Twelve rounds of the Decoy-SELEX process were performed to obtain a MRE specific for the pesticide malathion (Figure B.1). A ssDNA library with a diversity of 10^{15} molecules was enriched for those that bound to malathion and not the negative targets of selection (Figure B.2). The library was designed within our laboratory and designated RMW.N34. The ssDNA library was commercially synthesized and consists of two 23 base constant regions for polymerase chain reaction (PCR) amplification that flank a 34 base random region (Eurofins MWG Operon; Huntsville, AL).

The malathion derivative malathion monocarboxylic acid (Figure B.2b) was covalently biotinylated using EDC-NHS chemistry. It was reacted with 1-ethyl-3-[3-dimethylaminopropyl]carbodiimide (EDC) (ThermoScientific; Rockford, IL) and then incubated with Amine-PEG₂-Biotin (Pierce; Rockford, IL) according to manufacturer's instructions. The biotinylated derivative was bound to streptavidin-coated magnetic beads (New England Biolabs; Ipswich, MA) and washed to provide the immobilized target (IT).

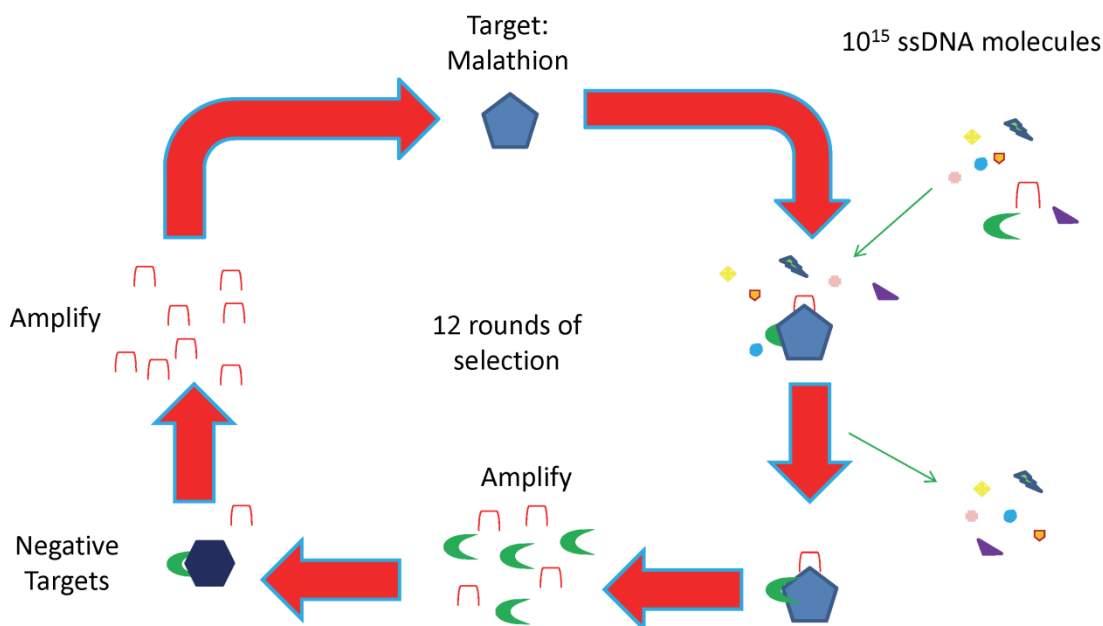


Figure B.1. Decoy-SELEX scheme for selection of a malathion MRE. 10¹⁵ different ssDNA molecules were incubated with malathion. Those that bound were amplified and subjected to closely-related targets. Those do not bind those were amplified and re-subjected to malathion under more stringent conditions. This process continued for 12 rounds.

For Round 1(+) selection, the ssDNA library was incubated with 50 μ L IT in a total of 500 μ L selection buffer containing 100 mM sodium chloride, 20 mM Tris-HCl, and 2 mM magnesium chloride (1X selection buffer; SB). The mixture was incubated at room temperature on a rotisserie for 48 hours. The IT with bound ssDNA was removed from the solution by magnetic separation, washed three times with 1 mL SB, and resuspended in SB. This was used

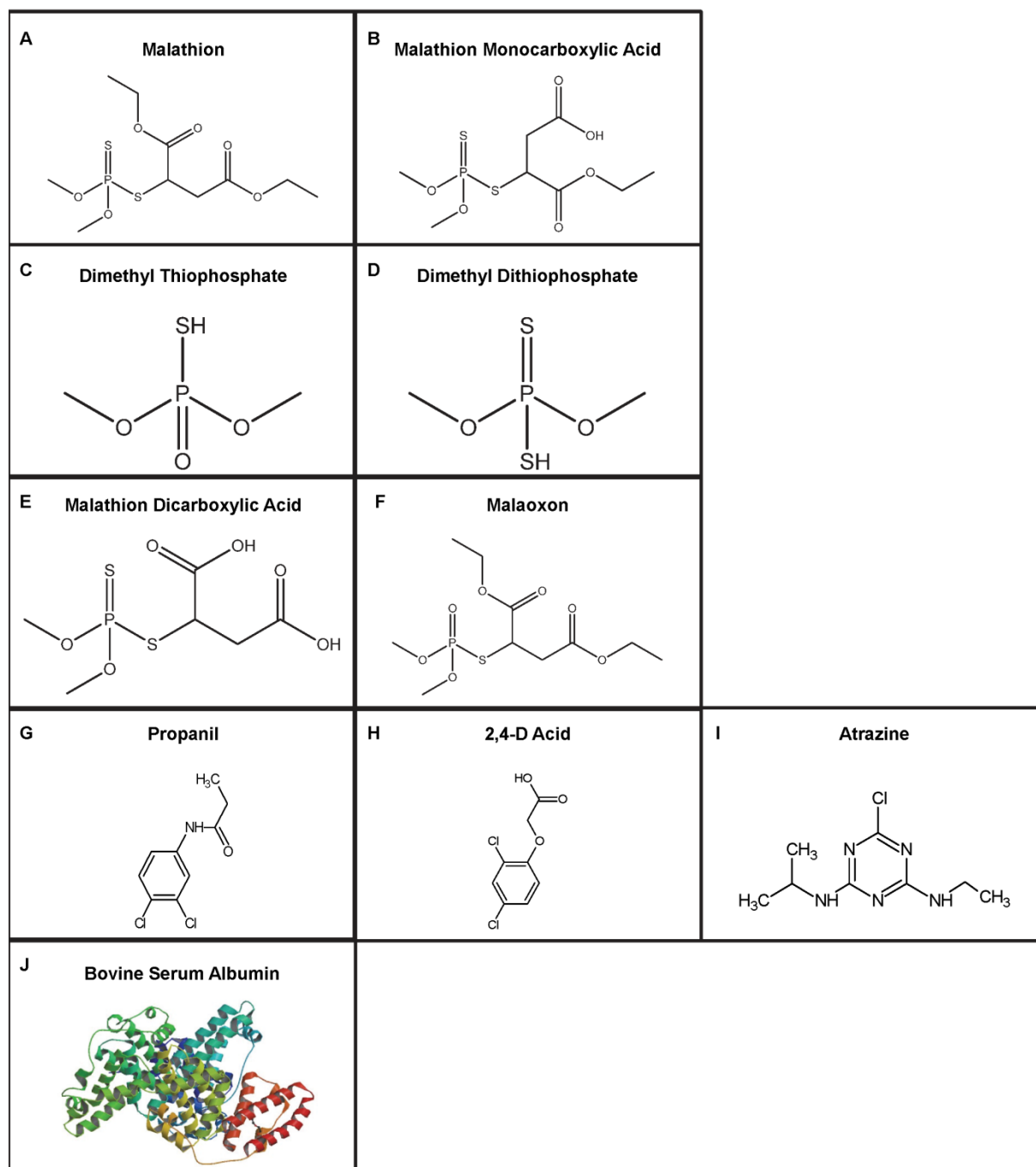


Figure B.2. Chemical structures of molecules used in selection of a malathion-specific MRE. (a) Structure of the target of selection, malathion. (b) Structure of malathion monocarboxylic acid, a malathion derivative used in magnetic bead immobilization. (c) and (d) Structures of dimethyl thiophosphate and dimethyl dithiophosphate used as negative targets in Round 8(-). (e) Chemical structure of malathion dicarboxylic acid, used in Round 9(-) selection. (f) Structure of malaoxon, used in Round 10(-) selection. (g), (h), and (i) Structures of propanil, 2,4-D acid, and atrazine, used successively in Round 7(-) of selection. (j) Ribbon structure of

bovine serum albumin (BSA) used as a negative target in Round 11(-) of selection (Bujacz 2012).

as a template for PCR amplification of ssDNA bound to the IT. PCR amplification was performed with the following reaction ingredients: enriched ssDNA library, 400 nM forward and biotinylated reverse RMW.N34 primers (Eurofins MWG Operon) (forward: 5'-TGTACCGTCTGAGCGATTCGTAC-3', biotinylated reverse: 5'-Biotin-GCACTCCTTAACACTGACTGGCT-3'), 250 μ M deoxynucleotide triphosphates, 1X GoTaq Reaction Buffer (Promega; Madison, WI), 3.5 units *Taq*, and pure water to 100 μ L. Thermal cycling conditions were: initial denaturation of 95°C for 5 minutes; cycling at 95°C for 1 minute, 63°C for 45 seconds, and 72°C for 1 minute; and final extension of 72°C for 7 minutes. Amplification for carrying the library to the next round of selection was performed using either a 4 mL or a 2 mL total volume PCR, which respectively preceded positive and negative selection rounds.

The amplified DNA was then purified by agarose gel electrophoresis using the GFX PCR purification kit (GE Healthcare; Piscataway, NJ). Recovered dsDNA was then incubated with streptavidin agarose resin (Pierce) for separation of single strands. The biotinylated reverse primer used in PCR amplification allowed for binding of the reverse strand to the agarose resin. This mixture was placed in a flow-through column and washed with five column volumes of phosphate-buffer saline (PBS). The single forward strand of amplified DNA was eluted from the column using five column volumes of 1 M sodium hydroxide. The elute ssDNA was prepared for ethanol precipitation by adding 0.1 volume of 3 M sodium acetate (pH 5.2), 2.5 volumes of cold 100% ethanol, and glycogen at 10 μ g/mL. The solution was frozen and then centrifuged at 13,000xg for 1 hour, the pellet washed with 70% ethanol, and centrifuged again for 15 minutes.

The pellet was dried under a vacuum and resuspended in SB. The amount of ssDNA present was verified to be at least 10^{13} sequences using a NanoDrop spectrometer (ThermoScientific).

Following Round 2(+) selection, the DNA was prepared for the next round as specified above. However, it was then incubated with immobilization substrate (IS) for Round 2(-) selection for 16 hours in 100 μ L total volume of SB. To prepare the IS, Amine-PEG₂-Biotin (Pierce) was incubated with streptavidin-coated magnetic beads (New England Biolabs) and washed to remove unbound biotin reagent. After the incubation period, the IS with bound ssDNA was separated from the solution by magnet. Washes with 25, 25, and 50 μ L of SB were performed and pooled with the total 100 μ L of incubation supernatant. This served as a template for PCR amplification, which was performed as previously described. The subtracted library was then prepared for Round 2(+) selection as described above.

Round 2(+) selection proceeded similar to Round 1(+), however the incubation period was 22 hours. Positive selections occurred in this manner, with decreasing incubation period for increased stringency, through Round 6(+). Negative selections similar to Round 1(-) were performed in Round 3 and 5, with longer incubation periods, to remove molecules binding to anything in the incubation that was not the malathion metabolite.

Beginning in Round 7(+), a competitive elution was performed in order to obtain a MRE which bound to malathion in solution. For this, the enriched library was incubated with 50 μ L of IT in a total volume of 500 μ L SB for 3 hours. The beads were then washed three times with 1 mL SB. However, in this round, 100 μ L of 10 mM malathion in SB was then added to the IT and incubated for 5 minutes. The IT was then removed from the solution by magnet, and the supernatant served as the template for PCR amplification. The enriched and amplified library was then prepared for Round 7(-) selection as noted above. This method of selection by

competitive elution was performed through Round 12 (+) with decreasing incubation times. Concentrations of malathion used for elution were also decreased, until finally in Round 12(+) the library incubation with the IT was immediate, and elution with 100 μ L of 100 nM malathion was immediate.

In Round 7(-), competitive elutions were performed with other pesticides found in similar environmental locations as malathion. The enriched library was incubated with the IT, the supernatant removed, and the beads washed 3 times. Then, the beads were incubated for 24 hours at room temperature on the rotisserie with 500 μ L of 1 μ M propanil in SB. The supernatant was removed and the beads were washed twice with 1 mL SB. The IT was then incubated with the same volume and concentration of 2,4-D Acid, washed, incubated with atrazine, and washed. The beads were then resuspended in SB and used as a template in PCR amplification. The DNA was then prepared for the next positive round of selection.

After Round 8(+) selection, Round 8(-) selection was done similar to the previous negative round, however dimethyl thiophosphate, then dimethyl dithiophosphate were used as negative targets. Following Round 9(+), Round 9(-) was performed with 500 μ L of 1 mM malathion dicarboxylic acid for 15 minutes. Round 10(-) selection was performed with 500 μ L of 1 mM malaoxon with an incubation period of 1 minute. The final negative selection, Round 11(-), was performed in the same manner with bovine serum albumin as the elutant.

B.2.2 Sequencing of the Enriched ssDNA Library

Following every third round of selection, a random sample of at least 30 ssDNA molecules present in the enriched library was taken. The library was PCR amplified with non-biotinylated primers as described above and ligated into the pCR2.1 vector (Invitrogen; Carlsbad, CA) according to manufacturer's instructions. This was cloned into competent bacteria, which

were plated onto selective agar. Colonies with inserts were amplified, and plasmids containing individual library inserts were purified using the AxyPrep Plasmid Prep Kit (Axygen; Union

Table B.1. Decoy-SELEX outline for identification of malathion MRE

Round	Positive Selection	Negative Selection
1	Immobilized Target (IT) 48 hrs	Immobilization Substrate (IS) 16 hrs
2	IT 22 hrs	-
3	IT 16 hrs	IS 24 hrs
4	IT 10 hrs	-
5	IT 6.5 hrs	IS 23 hrs
6	IT 3 hrs	-
7	IT 3 hrs, 5 min Competitive Elution (CE) w/ 10 mM Malathion (100 uL)	IT 3 hrs, 24 hr CE w/ 1 μ M Propanil, 2,4-D Acid, Atrazine in succession (500 uL)
8	IT 1 hr, 5 min CE w/ 1 mM Mal (100 uL)	IT 1 hr, 24 hr CE w/ 1 μ M dimethyl thiophosphate, dimethyl dithiophosphate in succession (500 uL)
9	IT 15 min, 5 min CE w/ 100 uM Mal (100 uL)	IT 15 min, 15 min CE w/ 1 mM Malathion Dicarboxylic Acid (500 uL)
10	IT 1 min, 1 min CE w/ 10 uM Mal (100 uL)	IT 1 min, 1 min CE w/ 1 mM Malaoxon (500 uL)
11	IT 1 min, immediate CE w/ 1 uM Mal (100 uL)	IT 1 min, 15 min CE w/ 1 mM BSA (500 uL)
12	IT immediate, immediate CE w/ 100 nM Mal (100 uL)	-

Immobilization target (IT) is malathion monocarboxylic acid bound to magnetic beads. Immobilization substrate (IS) is streptavidin-coated magnetic beads with biotin reagent. Competitive elution (CE) is removal of bound ssDNA from IT with target or negative target in solution. Times listed are incubation periods in hours (hrs) or minutes (min).

City, CA). Purified plasmids were sent for sequencing (Eurofins MWG Operon) using the

M13R primer complimentary to a region upstream of the pCR2.1 vector. At least 30 sequences

were obtained for each enriched library. These were then analyzed for consensus sequence families to identify structures binding to malathion.

B.2.3 Determination of MRE Binding Affinity

The Round 12 sequences were analyzed to produce consensus sequence families, and from those three were chosen for further characterization. These were chosen based on their inclusion in consensus families as well as their structure and stability as predicted by the Mfold DNA web server using buffer salt and 25°C as binding conditions (SantaLucia 1998; Peyret 2000; Zuker 2003). The chosen sequences were designated R12.20, R12.14, and R12.29. These sequences were subjected to fluorescent saturation binding assays to determine their affinity for malathion.

Concentrations of 0, 0.25, 0.5, 0.75, 1, 5, 10, 15, 20, and 25 nM MRE were used in saturation binding studies. In a 200 μ L total volume of SB, 10 μ L of IT was incubated with each concentration for five minutes. Unbound MRE was removed and each incubation was washed five times with 200 μ L SB, then the immobilized target was resuspended in 200 μ L SB and heated to 95°C for 10 minutes to denature and elute bound MRE. Eluted ssDNA in SB was placed in a 96-well microplate and measured in a Synergy 2 microplate reader equipped with a tungsten halogen lamp with excitation wavelength of 490 nm and an emission filter at 520 nm using Gen5 1.06 software (Biotek US; Winooski, VT). All fluorescence readings on the plate were normalized to 100 μ L of a 1 nM solution in SB of the fluorescent MRE and the no ssDNA incubation. For each concentration set, the same was done with washed streptavidin-coated magnetic beads to ensure binding over background to the beads. Each set of incubations were performed in triplicate. To determine the dissociation constant (K_d) of the MRE, data were analyzed with Origin 8 (OriginLab Corporation; Northampton, MA) using nonlinear regression

analysis and fit with the equation, $Y = ((B_{\max} * X) / (K_d + X)) + NS * X$, where B_{\max} is the maximum binding, K_d is the dissociation constant, and NS is nonspecific binding as previously described (Langan et al. 2012).

To determine binding of the selected MRE, R12.20, to compounds used in the selection, 50 nM fluorescent ssDNA was used. For each elutant used, in a 200 μ L total volume of SB, 10 μ L IT was incubated with R12.20 for five minutes. The magnetic beads were washed five times with 200 μ L SB. Then, each incubation was resuspended in 200 μ L of 1 μ M in SB of the following in SB: malathion, malathion monocarboxylic acid, dimethyl thiophosphate, dimethyl dithiophosphate, malathion dicarboxylic acid, malaoxon, propanil, 2,4, D acid, atrazine, and BSA. Each elutant was incubated with the immobilized target for ten minutes, and the solution was removed from the beads. This solution was placed in a 96-well plate and measured in a fluorescence plate reader. Data were normalized to an internal fluorescent standard of 1 nMMRE in 100 μ L SB and background fluorescence of each elution buffer. Each set of binding studies were performed in triplicate. Data were averaged and standard deviations calculated. For each elutant, a one-tailed t-test was performed to determine statistical differences in the means.

B.3 Results and Discussion

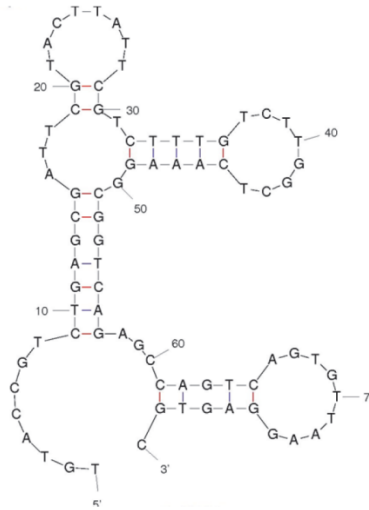
B.3.1 Identification of a Malathion-Specific MRE

In order to identify a MRE with high affinity for malathion that did not bind to other pesticides or closely-related structures, twelve rounds of Decoy-SELEX were completed (Table B.1). The selection ensured that the final MRE would not bind to the immobilization substrate, propanil, 2,4-D acid, atrazine, dimethyl thiophosphate, dimethyl dithiophosphate, malathion

A

R12.14	TGTACCGTCTGAGCGATTCTGACTATTCTGCTTTGTCTTGGCTCAAAGGCGGTCAAGCCAGTCAGTGTTAAGGAGTGC
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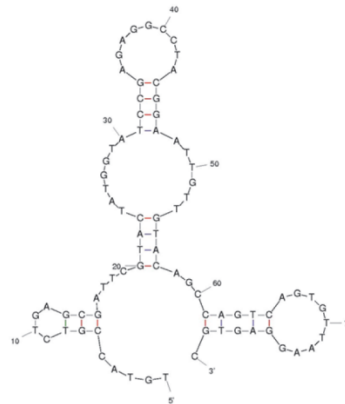
R12.13 TGTACCGTCTGAGCGATTCTGACTAAAAATAACCGCCGAGGTATCTCAGAGCGCGAGCCAGTCAGTGTTAAGGAGTGC
R12.14 TGTACCGTCTGAGCGATTCTGACTATTCTGCTTTGTCTTGG-CTCAAAGGCGGTCAAGCCAGTCAGTGTTAAGGAGTGC
R12.25 TGTACCGTCTGAGCGATTCTGACTACCGGTCAAGGAAAGG-TATCGCGACAGTCAGAGAGCCAGTCAGTGTTAAGGAGTGC



B

R12.20	TGTACCGTCTGAGCGATTCTGACTATGGTATCCGAGAGGCCCTACGGAATTGTTGTACAGCCAGTCAGTGTTAAGGAGTGC
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R12.20 TGTACCGTCTGAGCGATTCTGACTATGGTATCCGAGAGGCCCTACGGAATTGTTGTACAGCCAGTCAGTGTTAAGGAGTGC
R12.28 TGTACCGTCTGAGCGATTCTGACTACCGATTGAATTGGAGAGAGCTACGTCCTGTCTATTAGCCAGTCAGTGTTAAGGAGTGC
R12.46 TGTACCGTCTGAGCGATTCTGACTTAGGGAATTTCGCGATCTTGTCCCTCCAAAGAAGCCAGTCAGTGTTAAGGAGTGC



C

R12.29	TGTACCGTCTGAGCGATTCTGACTGTGCAGTGGCCTCGGACTAGAGTCTCTTCGTCAGCCAGTCAGTGTTAAGGAGTGC
--------	---

R12.18 TACCGTCTGAGCGATTCTGACTATATAGCCAGGTAGTATAGTCGGCCTAAAGAGAGCCAGTCAGTGTTAAGGAGTGC
R12.20 TGTACCGTCTGAGCGATTCTGACTATGGTATCCGAGAGGCCCTACGGAATTGTTGTACAGCCAGTCAGTGTTAAGGAGTGC
R12.29 TGTACCGTCTGAGCGATTCTGACTGTGCAGTGGCCT-CGGACTAGAGTCTCTTCGTCAGCCAGTCAGTGTTAAGGAGTGC

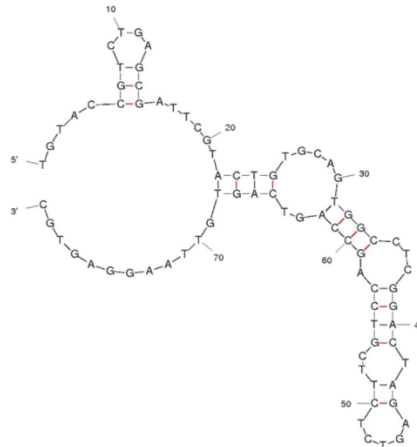


Figure B.3. Sequences of malathion MREs chosen from Round 12 library. For each, the sequence is above an example of a representative consensus sequence family of which the sequence is a member, and the structure as predicted by Mfold is on the bottom (SantaLucia 1998; Peyret 2000; Zuker 2003). Sequence and structure of (a) R12.14, (b) R12.20, and (c) R12.29.

dicarboxylic acid, malaoxon, or bovine serum albumin. Following Rounds 3, 6, 9, and 12, at least 30 random sequences were chosen from the enriched library for consensus sequence analysis. Sequences were analyzed and tracked throughout the *in vitro* selection. Three sequences were chosen, R12.14, R12.20, and R12.29 based on their presence in consensus families, structure, and stability as predicted by the Mfold DNA web server (Figure B.3). These sequences have a predicted Gibb's free energy value of -8.84 kcal/mol, -5.84 kcal/mol, and -7.89 kcal/mol, respectively.

B.3.2 Affinity and Specificity of Malathion-Specific MRE

Fluorescence saturation binding assays identified R12.20 as having the highest affinity and specificity (Figure B.3b). After assaying concentrations in the pM to low nM range, the dissociation constant (K_d) of this MRE was determined to be between 1.14 nM and 3.21 nM (Figure B.4). This is in the lower range of previously-selected ssDNA MREs for small-molecule targets (Hamula et al. 2006; Stoltenburg et al. 2007). This high affinity is likely to be a product of the increasingly stringent incubation conditions used in the Decoy-SELEX process.

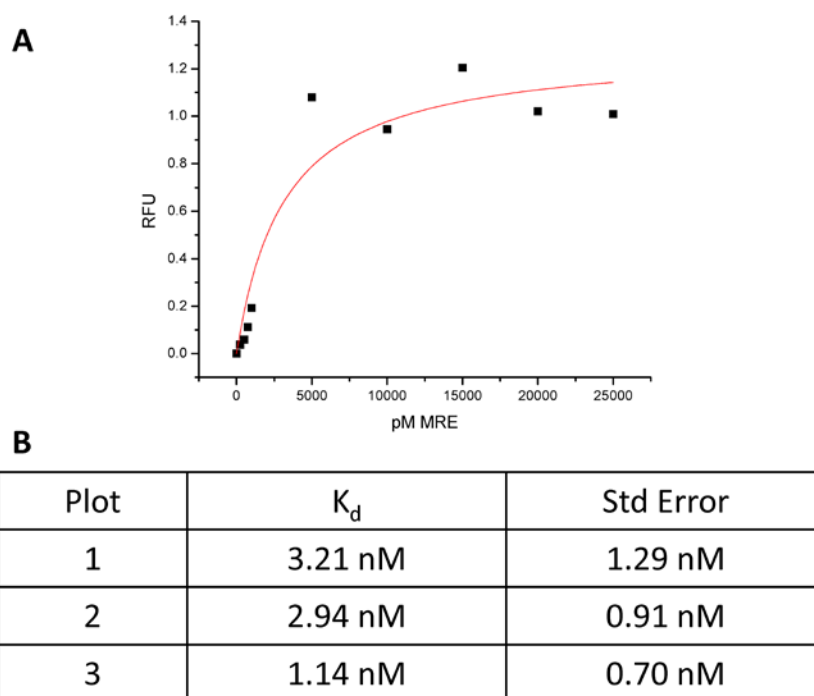


Figure B.4. Fluorescence equilibrium binding assays of R12.20 for malathion. (a) Representative saturation binding curve of R12.23 with nonlinear regression best fit. (b) Equilibrium dissociation constants (K_d) with standard error of three R12.23 binding assays.

The cross-reactivity of R12.20 for negative targets used in the selection was also determined. The ssDNA MRE binds to malathion significantly greater than it does to DMTP, DMDTP, malaoxon, propanil, and 2,4-D acid ($p < 0.05$) (Fig. 5). The binding of R12.20 to malathion is much higher than it is to atrazine (100% compared to 49%). Large variation, however, render this result statistically insignificant ($p = 0.06$). Binding to malathion monocarboxylic acid (MMA) is 70% that of malathion, however there is no difference in these samples ($p = 0.06$). This is expected as MMA was the molecule used to create the immobilized target used in all rounds of selection. Binding of the MRE to malathion dicarboxylic acid (MDA) is 60% that to malathion, however this is not significantly different ($p = 0.11$). The similarity of MMA and MDA explain the binding of R12.20 to MDA (Fig. 2b and 2e). Binding of the MRE is also as strong to BSA as it is to malathion. This may be due to the large, globular

nature of the protein and its interaction with ssDNA. This therefore suggests that R12.20 would be most successful as a sensing element in the presence of mostly small molecules. Additionally, even though binding to malathion is approximately two-fold higher than it to DMTP for example, the dissociation constant may be much higher than twice that of R12.20 for malathion. Therefore, incorporation into a sensor may lead to much greater specificity than it appears here.

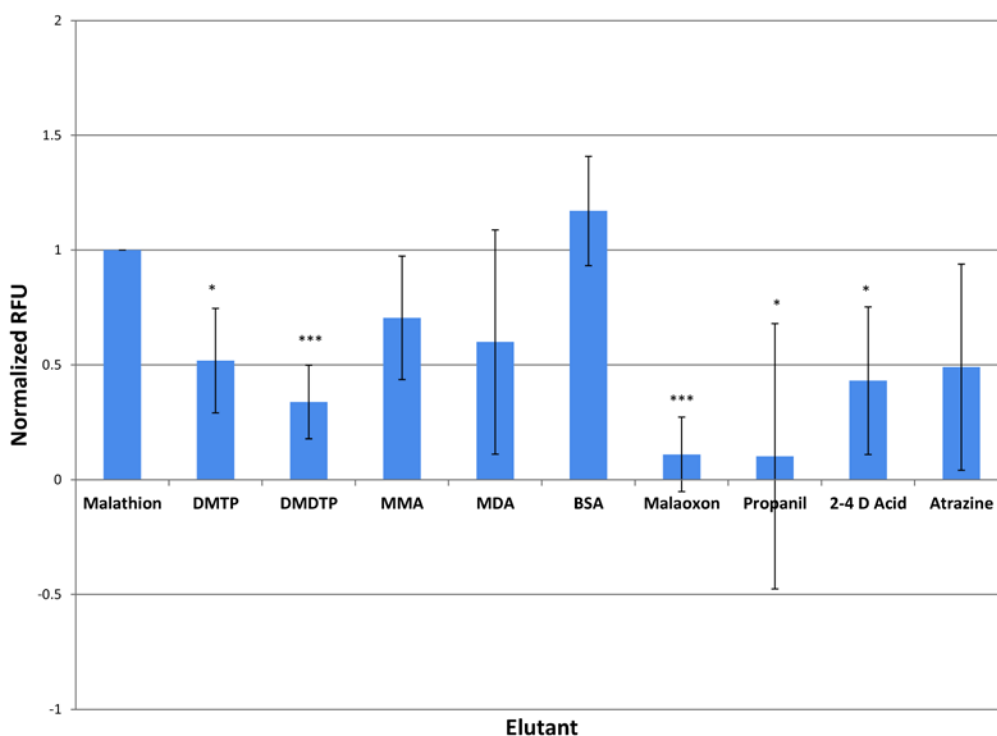


Figure B.5. Cross-binding studies of malathion MRE to other molecules used in the selection. Data were normalized to atrazine binding which was set to 100%, with error bars representing standard deviations. Statistical significance levels of $p < 0.05$ are designated by ‘*’, while levels of $p < 0.01$ are designated by ‘**’, and levels of $p < 0.001$ are designated by ‘***’. Malathion is the target of the selection and MMA was used to create immobilized target. DMTP, DMDTP, MDA, and malaoxon are molecules similar to malathion used as negative targets. BSA was also used as a negative target as model of a large, globular protein. Propanil, 2,4-D acid, and atrazine are other pesticides which were used as negative targets. MMA = malathion monocarboxylic acid; DMTP = dimethyl thiophosphate; DMDTP = dimethyl dithiophosphate; MDA = malathion dicarboxylic acid; BSA = bovine serum albumin.

The cross-binding results are particularly interesting as the MRE differentiates malathion from three closely-related structures, including malaoxon, that some antibodies cannot discern between (Yan et al. 2009). It is expected that even though the previously-selected antibodies were not tested for binding affinity or for binding to some of the metabolites tested here, the MRE will have higher affinity and specificity than antibodies previously identified for malathion (Nishi et al. 2003; Brun et al. 2005; Vera-Avila et al. 2005). This is because the *in vitro* selection process allows the investigator more control over final binding parameters than antibody selection. Furthermore, the Decoy-SELEX process allows the investigator greater control over which closely-related molecules the MRE will bind to.

The selected malathion ssDNA MRE will be useful for environmental sensing. Previously-developed nucleic acid MREs have been used as the sensing element in this capacity (Strehlitz et al. 2012; Yildirim et al. 2012). It is possible that design of the sensor will allow for a rapid, field-usable malathion detection device that will be quicker, cheaper, and easier to operate than current mechanisms. With easier detection of malathion, a greater sampling of contaminated environments will be possible. It is then possible to actively remediate these contaminated environments by biological, chemical, or physical methods (Nair and Pradeep 2007; Xie et al. 2009; Singhal et al. 2012).

B.4 Conclusions

Decoy-SELEX, a novel variation of *in vitro* selection, has been utilized to obtain a MRE for the pesticide malathion. Twelve rounds of selection were completed, with consensus sequence families identified and potential MREs obtained. The selected ssDNA MRE, R12.20, has a very high affinity malathion and selectivity compared to various other pesticides and

metabolites of malathion. This MRE will allow for rapid, cheap, and specific detection of malathion, which will lead to potential remediation of contaminated environments.

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Education

- Graduate
 - West Virginia University August 2008-Present
 - Ph.D. Student, Pharmaceutical and Pharmacological Sciences
 - ❖ August 2010-Present
 - ❖ Ph.D. Candidate, August 2011
 - Ph.D. Student, Department of Biology
 - ❖ August 2008-July 2010
- Undergraduate
 - University of Virginia August 2004-May 2008
 - B.A., Biology
- High School
 - Graham High School, Bluefield, Va. August 2000-May 2004
 - Advanced Degree, Summa Cum Laude

Research Experience

- Graduate Research Assistant
 - Dr. Letha Sooter January 2010-Present
 - Investigations in Molecular Recognition Element Selection and Single-Wall Carbon Nanotube Properties
 - Dr. Rajesh Naz August 2008-December 2009
 - Novel Biomarkers and Immunotherapies for Prostate Cancer
- Undergraduate Research Assistant
 - Dr. Edmund Brodie III September 2006-May 2008
 - Evolutionary Genetics of Caribbean *Anolis* spp.

Fellowships and Assistantships

- American Foundation for Pharmaceutical Education (AFPE) Pre-Doctoral Fellowship in the Pharmaceutical Sciences
September 1, 2011-August 31, 2013

- WVNano/NanoSAFE Graduate Fellowship Program (NSF Cooperative Agreement)
August 15, 2011-August 15, 2013
- Cancer Nanotechnology Graduate STEM Fellowship
Selected for period of August-December 2011
- Graduate Research Assistant, West Virginia University
 - Dept. of Basic Pharmaceutical Sciences
August 2010-Present
 - Dept. of Biology
May 2010-August 2010
 - Dept. of Microbiology and Immunology
May 2009-August 2009
- Graduate Teaching Assistant, West Virginia University Dept. of Biology
 - Biol 321: The Total Science Experience
Spring 2009, 2010
 - Biol 441: Vertebrate Microanatomy
Fall 2009
 - Biol 115: Intro Biology Lab
Fall 2008

Publications and Patents

Williams RM, Hajiran CJ, Nayeem S, and Sooter LJ. Identification of an antibody fragment molecular recognition element specific for prostate cancer cells. In Preparation for *Molecular Oncology*.

Williams RM, Eamonn Maher, and Sooter LJ. *In vitro* selection of ssDNA molecular recognition elements for the pesticide malathion. Submitted to *Applied Biochemistry and Biotechnology*, March 2013.

Williams RM and Sooter LJ. *In vitro* selection of molecular recognition elements specific for the herbicide atrazine. Submitted to *PLoS One*, March 2013.

Williams RM, Taylor H, Thomas J, Hines B, and Sooter LJ. The effect of DNA- and sodium cholate-dispersed single wall carbon nanotubes on the green algae *Chlamydomonas reinhardtii*. Submitted to *Environmental Toxicology*, March 2013.

Williams RM, Nayeem S, Hines B, and Sooter LJ. The effect of DNA-dispersed single-wall carbon nanotubes on the polymerase chain reaction. Under Review for *Nanotechnology*. Submitted December 2012.

Williams RM and Sooter LJ. *In vitro* selection of cancer cell-specific molecular recognition elements from amino acid libraries. In Preparation for *Molecular Cancer Research*.

Williams RM and Naz RK. (2010) Novel biomarkers and therapeutic targets for prostate cancer. *Frontiers in Bioscience* S2:677-684.

Letha J. Sooter and Ryan M. Williams. Single-stranded DNA sequences that bind atrazine and simazine. Disclosure of Intellectual Property, West Virginia University Docket No. 00584. Submitted December 2011.

Letha J. Sooter, Kailey Imlay, and Ryan M. Williams. Single-stranded DNA sequences that bind to *Pseudomonas aeruginosa* Exotoxin A. Disclosure of Intellectual Property, West Virginia University Docket No. 00585. Submitted December 2011.

Awards and Honors

- Student Travel Award, American Institute of Chemical Engineers Society for Biological Engineering (AIChE SBE) 2012 International Conference on Bioengineering and Nanotechnology. Berkeley, CA. June, 2012
- Rho Chi National Pharmacy Honor Society, Alpha Mu Chapter April, 2012-Present
- George M. Cruise Foundation Scholarship, First Community Bank, Bluefield, WV 2004-2012
- Ethel N. Bowen Foundation Scholarship, First Century Bank, Bluefield, WV 2004-2012
- Cleo Lawson Mitchell Foundation Scholarship, Graham High School, Bluefield, Va. 2004-2008
- Lions Club of Bluefield, Va Scholarship 2004
- Madhu Chopra Memorial Foundation, Bluefield, Va. 2004
- Graham High School Women's Boosters Foundation Scholarship, Bluefield, Va. 2004
- Peacemaker's Scholarship Foundation, Abingdon, Va. 2004
- Gift of Hope 21st Century Scholars Program, United Methodist Scholarship Fund, Nashville, TN 2004

Laboratory Mentoring Experience

- 1 High School Student
- 15 Undergraduate Students
- 10 Graduate Rotation Students

- 4 Graduate Students
- 1 Pharm. D. Student

Oral and Poster Presentations-External

Williams RM, Hajiran CJ, Nayeem S, and Sooter LJ. In Vitro Selection of Molecular Recognition Elements for Molecular Targeting of Prostate Cancer. EORTC-NCI-AACR Symposium on Molecular Targets and Cancer Therapeutics. Dublin, Ireland. Abstract # 508. November 2012.

Sooter LJ and Williams RM. Identification and Targeting of Prostate Cancer Cells Via Molecular Recognition Elements. American Institute of Chemical Engineers Society for Biological Engineering (AIChE SBE) 2012 International Conference on Bioengineering and Nanotechnology. Berkeley, CA. June 2012.

Williams RM and Sooter LJ. In vitro selection of DNA molecular recognition elements for the pesticide atrazine. West Virginia Academy of Sciences STaR Symposium, Institute, WV, **Oral Presentation**. April, 2012.

Williams RM, Nayeem S, and Sooter LJ. *In vitro* selection of prostate cancer cell-specific molecular recognition elements. American Association of Cancer Researchers (AACR) 2012 Annual Meeting, Chicago, IL. Abstract # 888. April, 2012.

Williams RM, Nayeem S, and Sooter LJ. Antibody fragments for the targeted treatment and diagnosis of prostate cancer. Appalachian Health Summit, University of Kentucky Center for Clinical and Translational Science. Lexington, KY. March, 2012.

Williams RM, Nayeem S, and Sooter LJ. *In Vitro* Selection of Prostate Cancer Cell-Specific Molecular Recognition Elements. American Association of Pharmaceutical Scientists (AAPS) Pittsburgh Chapters, Student Research Symposium, Duquesne University. October 2011.

Oral and Poster Presentations-Internal

Williams RM, Nayeem S, and Sooter LJ. Identification of Antibody Fragments Specific for Prostate Cancer Cells *Via* SELEX. West Virginia University Linking Innovation, Industry, and Commercialization (LIINC) Biomedical Sciences Graduate Student Event, **Invited Oral and Poster Presentation**. March 2013.

Williams RM, Nayeem S, and Sooter LJ *In vitro* selection of antibody fragments that specifically bind to prostate cancer cells. West Virginia University Health Sciences Center Van Liere Research Day. March 2012.

Williams RM, Nayeem S, and Sooter LJ. Molecular Recognition Elements for Prostate Cancer and Pesticides. West Virginia University Linking Innovation, Industry, and Commercialization (LIINC) Biomedical Sciences Graduate Student Event, **Invited Oral and Poster Presentation**. February 2012.

Williams RM, Nayeem S, and Sooter LJ. *In Vitro* Selection of Prostate Cancer Cell and Pesticide Molecular Recognition Elements. West Virginia University School of Pharmacy Research Day. October 2011.

Williams RM and Sooter LJ. Systematic Evolution of Ligands for Prostate Cancer Cells and Atrazine by Exponential Enrichment. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.

Williams RM and Sooter LJ. “Molecular Recognition Elements for Prostate Cancer Cells and Atrazine.” WVNano Brown Bag Lunch Speaker Series **Oral Presentation**. November 2010.

Williams RM and Sooter LJ. *In Vitro* Selection of Prostate Cancer Specific and Herbicide Binding Elements. West Virginia University School of Pharmacy Research Day. October 2010.

Williams RM and Sooter LJ. Prostate Cancer Detection by Molecular Recognition Elements. West Virginia Academy of Sciences Symposium. April 2010.

Williams RM and Sooter LJ. Prostate Cancer Detection by Molecular Recognition Elements and Environmental Toxicity of Single-Walled Carbon Nanotubes. West Virginia Nano Symposium. April 2010.

Williams RM and Sooter LJ. Quantum Dot-Conjugated Molecular Recognition Elements in Detecting Prostate Cancer. West Virginia University American Chemical Society Symposium. March 2010.

Other Presentations

Hajiran CJ, Williams RM, and Sooter LJ. Systematic evolution of ligands for prostate cancer cells and bromacil by exponential enrichment. NanoSAFE SURE Symposium. July 2012.

Jackson AM, Williams RM, and Sooter LJ. *In vitro* selection of molecular recognition elements for detection of Exotoxin A and prostate cancer. NanoSAFE REU Symposium. July 2012.

Kulick AR, Williams RM, and Sooter LJ. *In Vitro* Selection of PETN Molecular Recognition Elements Using Capillary Electrophoresis. West Virginia Academy of Sciences STaR Symposium, Institute, WV. April, 2012.

Imlay K, Williams RM, and Sooter LJ. Isolation of molecular recognition elements against Exotoxin A for application in toxin detection. West Virginia Academy of Sciences STaR Symposium, Institute, WV. April, 2012.

Zimmermann N, Kulick AR, Williams RM, and Sooter LJ. Selections of odor binding Molecular Recognition Elements using CE-SELEX. West Virginia Academy of Sciences STaR Symposium, Institute, WV. April, 2012.

Imlay K, Williams RM, and Sooter LJ. Selection of Molecules that Bind Exotoxin A for Application in Toxin Detection. West Virginia University Health Sciences Center Van Liere Research Day. March 2012.

Kulick AR, Williams RM, and Sooter LJ. *In Vitro* Selection of PETN Molecular Recognition Elements Using Capillary Electrophoresis. West Virginia University Health Sciences Center Van Liere Research Day. March 2012.

Zimmermann N, Kulick AR, Williams RM, Daly K, and Sooter LJ. *In vitro* selection of odor-specific aptamers using CE-SELEX. West Virginia University Health Sciences Center Van Liere Research Day. March 2012.

Imlay K, Williams RM, and Sooter LJ. Isolation of Molecules That Bind Exotoxin A for Application in Toxin Detection. Undergraduate Research Day at the Capitol, Charleston, WV. January 2012.

Kulick AR, Williams RM, and Sooter LJ. Identification of Molecules that Bind to the Explosive PETN. Undergraduate Research Day at the Capitol, Charleston, WV. January 2012.

Imlay K, Williams RM, and Sooter LJ. Molecular Recognition Elements against Exotoxin A for Applications in Toxin Detection. West Virginia University School of Pharmacy Research Day. October 2011.

Kulick AR, Williams RM, and Sooter LJ. *In Vitro* Selection of PETN Molecular Recognition Elements Using Capillary Electrophoresis. West Virginia University School of Pharmacy Research Day. October 2011.

- Sooter LJ, Williams RM, Taylor H, Nayeem S, and Dolash BD. Effects of Single-Walled Carbon Nanotubes on Green Algae and the Polymerase Chain Reaction. West Virginia University School of Pharmacy Research Day. October 2011.
- Yedlapalli SL, Williams RM, Holland LM, and Sooter LJ. Two Molecular Recognition Element Selections Against Environmental Pollutants. West Virginia University School of Pharmacy Research Day. October 2011.
- Zimmermann N, Kulick AR, Williams RM, Daly K, and Sooter LJ. Identification of Odorant-Specific Aptamers Using CE-SELEX. West Virginia University School of Pharmacy Research Day. October 2011.
- Imlay K, Williams RM, and Sooter LJ. *In Vitro* Selection of Molecular Recognition Elements Against Exotoxin A. WVNano SURE Symposium. July 2011.
- DeBerry SM, Williams RM, and Sooter LJ. *In Vitro* Selection for DNA Aptamers Targeting PETN. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.
- Douglas IP, Williams RM, and Sooter LJ. Selecting Molecular Recognition Elements Against HGPIN Cells Using a Yeast Displayed scFv Library. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.
- Ghabra O, Williams RM, and Sooter LJ. Forensic Identification Via Hand Bacterial Community Analysis. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.
- Imlay K, Williams RM, and Sooter LJ. *In Vitro* Selection of Molecular Recognition Elements Against Exotoxin A. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.
- Rhodes GT, Williams RM, and Sooter LJ. A Molecular Recognition Element for the Detection of O,O-Diethyl Dithiophosphate. West Virginia University Health Sciences Center Van Liere Research Day. March 2011.
- Taylor H, Williams RM, and Sooter LJ. The Effects of DNA-dispersed and Sodium Cholate-dispersed Single-Walled Carbon Nanotubes on Algae *Chlamydomonas reinhardtii*. West Virginia University Health Sciences Center Van Liere Research Day. March 2011. First place-undergraduate poster competition.
- Douglas I, Williams RM, and Sooter LJ. Isolation of Antibody scFv Specific to PIN cells. West Virginia University School of Pharmacy Research Day. October 2010.

Thomas J, Williams RM, and Sooter LJ. Toxicity of single-walled carbon nanotubes to algae.
West Virginia University School of Pharmacy Research Day. October 2010.

Activities

- Rho Chi National Pharmacy Honor Society, Alpha Mu Chapter 2012-Present
- Member American Association of Cancer Researchers (AACR) 2011-Present
- Member American Association of Pharmaceutical Scientists (AAPS)
 - West Virginia University Student Chapter 2011-Present
- Member American Chemical Society (ACS) 2011-2012
- WVU Cell Biology Training Program 2010-2012
- WVNano NanoDays Volunteer Community Outreach March 17, 2012
- National Science Foundation Workshop, Science: Becoming the Messenger, West Virginia University, Morgantown, WV October 11, 2011
- WVU Biology Graduate Student Association Treasurer 2009-2010
- Cavalier Daily Newspaper Associate Sports Editor and Staff Writer 2007-2008
- University of Virginia Children's Hospital Pediatric Unit
 - Over 60 volunteer hours 2007-2008
- Energy Express (AmeriCorps) Mentor, Mercer County, WV
 - Over 300 volunteer hours per summer (>900 hours total) Summers 2005, 2006, 2007
- Martha Jefferson Hospital Emergency Department, Charlottesville, Va.
 - Over 60 volunteer hours 2006-2007
- University of Virginia Madison House Boosters Program
 - Over 30 elementary classroom volunteer hours 2005-2006

Investigations in Molecular Recognition Element Selection and Single-Wall Carbon Nanotube Properties

Ryan Martin Williams


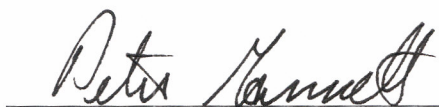
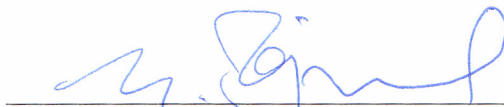

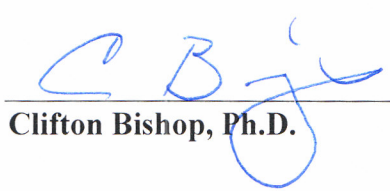
A dissertation submitted to the School of Pharmacy at
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degree of

Doctor of Philosophy in Pharmaceutical and Pharmacological Sciences

Department of Basic Pharmaceutical Sciences

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2013

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 Peter Gannett, Ph.D.	3/4/12 Date
 Yon Rojanasakul, Ph.D.	3/4/12 Date
 Linda Vona-Davis, Ph.D.	3/4/2013 Date
 Clifton Bishop, Ph.D.	3/4/13 Date